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AUTOLOGOUS IMMUNE CELL THERAPY: CELL COMPOSITIONS. METHODS AND APPLICATIONS TO TREATMENT OF HUMAN DISEASE RELATED APPLICATIONS

This application is a continuation-in-part of U.S. application Serial No. 08/506,668, filed July 25, 1995, by Micheal Gruenberg, entitled PROCESS FOR PRODUCING EFFECTOR IMMUNE CELLS FOR USE IN ADOPTIVE CELLULAR IMMUNOTHERAPY. Benefit of priority thereto is and the state

This application is also related to U.S. application Serial No. 08/506,173, filed July 25, 1995, by Micheal Gruenberg, entitled CELL GROWING DEVICE FOR IN VITRO CELL POPULATION EXPANSION

10 For purposes of the U.S. national stage, the subject matter of each of U.S. application Serial Nos. 08/506,668 and 08/506,173 is herein incorporated by reference its entirety.

FIELD OF INVENTION

herein claimed 19 19 f

This invention is directed to methods of adoptive immunotherapy. 15 In particular, methods of autologous cell therapy are provided. Compositions containing substantially homogeneous populations of functionally or phenotypically defined immune cells that have been isolated from a patient, differentiated and/or expanded ex vivo are provided. Uses of such compositions for treating or preventing disease or 20 otherwise altering the immune status of the patient by reinfusing such cells are also provided.

BACKGROUND OF INVENTION

T lymphocytes are immune cells that are primarily responsible for protection against intracellular pathogens and suppression or elimination 25 of certain tumors. Mature T lymphocytes, which all express the CD3 cell surface antigen, are subdivided into two subtypes, based on expression

of either the CD4 or CD8 surface antigen. CD4* T cells recognize antigen presented in association with class II major histocompatibility complex (MHC) molecules. CD4* cells are generally involved in regulatory functions in immune responses by virtue of the cytokines they produce. These cytokines, such as IL-2, mediate an immune cell attack on a pathogen or an antibody attack against an inveding organism.

CD8* T cells recognize antigen presented in association with class I MHC molecules. CD8* cells are involved in effector functions in immune responses, such as cytotoxic destruction of cells bearing foreign antigens. The cells that mediate these responses are designated cytotoxic T lymphocytes (CTLs). These cells, which are generally CD8* cells (although some are CD4*) represent a mechanism for resistance to viral infections and tumors. The effector function of CTLs is dependent upon the cytokine production from CD4* regulatory cells.

Adoptive immunotherapy is an experimental treatment method designed to boost a patient's immune response against a virus or a tumor. The method involves the removal of immune cells from an individual, the forming of effector cells outside the body (ex vivo), the expansion of the cells to clinically-relevant numbers and the re-infusion of the cells into the patient. Adoptive immunotherapy protocols have not been made commercially available and are not in widespread use because of the extreme toxicities associated with the infusion of the interleukin-2 (IL-2) with the cells. IL-2 is used in these protocols to cause the differentiation and/or expansion of effector immune cells. Immune cells cultivated in IL-2, however, become dependent on the cytokine for continued viability and effector function, thus necessitating the infusion of IL-2 together with the effector cells. All adoptive immunotherapy protocols

involving differentiated effector cells incorporate the use of IL-2.

The severe toxicity associated with the use of IL-2 has limited the application of adoptive immunotherapy to the treatment of terminally-ill cancer patients and the treatment of viral infections in AIDS patients.

Adoptive immunotherapy and the use thereof for treating cancer.

5 The first attempts at adoptive immunotherapy in humans employed lymphokine activated killer (LAK) cells, which are immune effector cells functionally defined by their ability to lyse fresh tumors. LAK cells are produced when peripheral blood mononuclear cells are exposed to high concentrations of IL-2 ex vivo [see, e.g., Grimm, et al. (1982) J. Exp. 10 Med. 155:1832]. To produce LAK cells for use in treating cancer patients [see, U.S. Patent No. 4,690,915], leukocytes are removed from a cancer patient and exposed to high levels of JL-2 for 3-6 days, which causes a portion of the cells to differentiate into LAK cells: The resulting heterogeneous population of cells is reinfused to the donor concomitant with a high systemic dose of IL-2. As noted, the high systemic doses of IL-2 are highly toxic and not well tolerated. Methods in which the potency of LAK cells is increased have been developed. It has been observed [see, e.g., U.S. Patent No. 4,849,329] that the addition of an L-amino acid with IL-2 during the ex vivo 20 differentiation step increases the LAK activity of the resulting cells 4-5 fold. Administration of LAK cells with IL-2 and an ornithine decarboxylase inhibitor enhances the effectiveness of the treatment (see, U.S. Patent No. 5,002,879]. Exposure of lymphocytes to an anti-CD3 monoclonal antibody (mAb) during the LAK differentiation stage of the process produces effector cells with enhanced anti-tumor activity [U.S. Patent No. 5,326,763), and use of IL-7, with or without IL-2, in the LAK differentiation step can also produce more potent LAK effector cells [see, U.S. Patent No. 5,229,115]. The administration of GM-CSF with IL-2

has also been reported to cause an increase in LAK activity [see.....

Takahashi, et al. (1995) <u>Jap. J. Cancer Res. 86</u>:861). All protocols, however, require administration of IL-2.

Early clinical results of adoptive immunotherapy using LAK cells in terminally-ill cancer patients; particularly thôse with manginer mannoma, 5 had reported response rates of 21:44% (see, e.d., Rosenberg et al. (1985) N.Engl. J. Med. 313:1485 and Rosenberg et al. (1987) N. Engl. J. Med. 316:889). Results of more recent phase il clinical studies, while still showing promise, have produced a broad range of response rates from 0-33% (see, e.g., Dillman, et al. (1991) J. Clin. Oncol. 9:1233.

10 Thompson, J.A. et al. (1992) J. Clin. Oncol. 10:960; Foon, et al. (1992) J. Immunother: 11:1984 and Roreks et al. (1991) Facts. Surg. 26:868). The differences in response rates are attributed, partly, to Variations in dosages of LAK cells and IL-2 administrated, and the differences in tumor-killing activities of the heterogeneous populations of CAK cells.

Methods for generating a relatively homogenous population of LAK cells for adoptive immunotherapy have been developed is e. U.S. Patent No. 5,057/423]. The process described in U.S. Patent No. 5,057/423]. The process described in U.S. Patent No. 5,057/423 population of LAK progenitor cells (LGL) from the peripheral blood mononuclear cells. These LGL are then exposed to IL-2, which causes a majority of the LGL to differentiate into LAK cells. The resulting effector cells, known as A-LAK, have been shown to be effective in killing human carcinoma in hude mice (see, Sacchi (1991) at al. int. U. Cancer 47:784) Boiardi, at al. (1994) Cancer Immunol. Immunoth, 39:1931. It is exceedingly difficult, however, to produce sufficient numbers of A-LAK from humans. Even with the use of teeder cells to improve as vivo expansion, A-LAK cultures from approximately 60% of cancer patients demonstrated inadequate expansion issee.

Sedimavr. et al. (1991) J. Immunother, 10:3361.

Another adoptive immunotherapy protocol involves the administration of autologous tumor infiltrating lymphocytes (TIL) to cancer patients. TIL cells are more potent at killing tumors than LAK cells in animal experiments, but are difficult and expensive to generate for treatment of patients. TIL cells are autologous effector cells differentiated in vivo in solid tumors [see, U.S. Patent No. 5,126,132, which describes a method for generating TIL cells for adoptive immunotherapy of cancer). TIL cells are produced by removing a tumor sample from a patient, isolating lymphocytes that were infiltrating into 10 the tumor sample, growing these TIL cells ex vivo in the presence of IL-2 and reinfusing the cells to the patient along with IL-2. A 60% response rate in evaluable cancer patients using this protocol has been reported. [see, Rosenberg, et al. (1988) N. Engl. J. Med. 319:1676]. Another study reported a 23% response rate [see, Dillman, et al. (1991) Cancer-15 68:1]. It, however, has been difficult to consistently propagate sufficient numbers of TIL cells for use in adoptive immunotherapy protocols. ... In addition, the type of immune cells derived from TIL cultures are extremely variable. The cells recovered from tumor samples contain pure or mixed populations of cells with differing activities and potencies. 20 Some cells are produced with MHC-restricted anti-tumor cytolytic activity, some with non-MHC restricted anti-tumor cytolytic activity and some without any anti-tumorcytolytic activity. Also, other than cultures derived from melanomas, cultures of TIL cells rarely produce tumorspecific cells from patients with solid tumors; and tumor-specific cells are produced only from about 50-75% of patients with metastatic melanoma. Because TIL cell therapy is associated with extreme toxicity

Because TIL cell therapy is associated with extreme toxicity associated with infusion of IL-2, efforts have been made to enhance the efficacy of the treatment. For example, addition of IL-10 with IL-2 has been shown to increase the anti-tumor function of TIL cells in mice [see.

Yang, et al. (1995) J. Immunol, 155:3897. Increasing the II-6 concentration at the tumor site has also been shown to result in enhanced anti-tumor activity in TIL cells from mice [see, Marcus, et al. (1994) J. Immunoth. Emphasis Tumor Immunol. 15:105). The anti-tumor activity of TIL cells is also increased by activating tumor draining lymph node cells with anti-CD3 mAb in the presence of IL-1 [see, Hammel, et al. (1994) J. Immunoth: Emphasis Tumor Immunol. 16:11.

Because of the variability in the effector function of cells derived from tumor infiltrates or draining lymph nodes, effort is being invested in 10 development of methods to promote the ex vivo sensitization of fumorreactive immune cells for use in adoptive immunotherapy of cancer. Tumor-antigen specific, MHC-restricted CTL from precursor cells present in the cellular infiltrates of breast cancer patients have been produced by incubating precursor cells with recombinant avipox MAGE-1 la marker 15 present on a class of tumors], causing the formation of MAGE-1 specific CTL [(MAGE-1 and other MAGE antigens are antigens expressed on tumor cells); see Toso, et al. (1996) Cancer Research 56:16; see also U.S. Patent No. 5,512,444]. Another ex vivo sensitization method for generating potent MHC-restricted CTL involves the incubation of peripheral blood mononuclear cells (PBMC) from melanoma patients with autologous, irradiated PBMC that have been pulsed with synthetic peptides of qp100, a melanoma-associated antigen [see, Salgaller, et al. figure Mary of America (1995) Cancer Research 55:49721.

An alternative to TIL cells in adoptive immunotherapy of cancer are 25 "ALT" cells. These cells are ex vivo activated peripheral blood lymphocytes with CTL activity. They are activated in an IL-2-containing supernatant derived from a previously prepared one-way mixed lymphocyte culture or by using cytokine-rich, autologous supernatant harvested from a previous lymphocyte culture stimulated with anti-CD3

mAb. Monthly infusions of ALT cells, combined with daily oral cimetidine to reduce tumor-associated suppressor activity), significantly prolongs survival and induces durable tumor responses in renal cell carcinoma and melanoma patients [see, Graham, et al. (1993) Semin: Urol., 11:27-and Gold, et al. (1996) J. Surg. Res. 59:279)

Gold, et al. (1996) J. Surgy Res. 59:279] Abstract a page 9350 and 501. 3 Other effector immune cells have been used or proposed for adoptive immunotherapy of cancer. For example, the PWM-AK cell has been proposed as a possible candidate for adoptive immunotherapy of cancer. These effector cells are pokeweed mitogen activated PBMC with 10 similar activity to LAK cells (see, Ohno, et al. (1994) Int. 3. 4 6 5 3 3 3. Immunopharm, 16:761]. Human activated macrophages (MAK) have also been proposed as effector cells in adoptive immunotherapy of cancer. The MAK cells are differentiated from the peripheral blood by activation with interferon-v (IFN-v) and have been shown to cause regression of 15. experimental tumors in animals, but have not shown a clear therapeutic response in humans (see, Bartholeyns et al. (1994) Anticancer Research 14:2673]. Activated natural killer cells (ANK) have also been proposed for use in adoptive immunotherapy of malignancies. ANK cells are prepared by panning of peripheral blood stem cells on CD5/CD8 coated 20 flasks yielding a population enriched for monocytes or NK precursors and then treating the cells with high concentrations of IL-2. A humanderived, MHC non-restricted CTL clone (TALL-104) has also shown promise for use in adoptive immunotherapy protocols for cancer treatment when used in conjunction with IL-12 [see, Cesano, et al. (1994) J. Clin. Invest. 94:1076]. Increasing interest in the use of MAK, ANK and other mononuclear phagocytes in adoptive immunotherapy protocols for treatment of cancer has led to the development of improved methods to reproducibly harvest large numbers of functional human

circulating blood monocytes by counterflow centrifugal elutriation (see, Faradiji, et al.: (1994) J.: Immunol. Methods 174:297).

An emerging adoptive immunotherapy strategy for treatment of cancer is to isolate and/or generate antigen presenting cells such as

5 dendritic cells from a patient s.blood/pulse the cells with tumor.

fragments or antigenic peptides and then reintroduce the cells to the patient [see; Grabbe; et al. (1995) Immunol. Today 16:117]. Methods for obtaining large numbers of dendritic cells from precursors in the blood got adults have been described [see; Romani; et al. (1994) 3. Exp. Med.

10 180:83 and Bernhard; et al. (1995) Cancer Res. 55:1099].

Another application of immune cell adoptive immunotherapy is the treatment of viral disease. Adoptive immunotherapy protocols using viral-specific CD8+, and CD4+ effector cells have been developed for the

15 treatment of infections with CMV/EBV and HIV [set/Riddell et al."(1995)

Ann. Rev. Immunol. 13:545; wan Lunzen/et/al. (1995) Adv. Expf Med.

Biol. 374:57; and Klimas, et al. (1994) AIDS 6:1073) These protocols involve purifying CD8.#..T-cells from the peripheral blood of AIDS 10:10 patients, expanding the cells with phytohemagglutinin and IL-2 and re-

20 Infusing the cells, with concomitant IL-2 infusion, to the patient [see, Whiteside, et al. (1993) <u>Blood 81</u>:2085; Klimas, et al. (1994) <u>AIDS</u> 8:1073; Riddell, et al. (1993) <u>Curr. Opin Immunol. 5</u>:484; Torpey, et al. (1993) <u>Cline Immunol. Immunopath.</u> 68:263; Ho, et al. (1993) <u>Blood 681</u>:2093, and Riddell, et al. (1992) <u>Science 257</u>:238].

25 Methods for growing immune cells in vitro

A majority of adoptive immunotherapy protocols are hampered by the inability to grow clinically relevant (he inability to grow clinically relevant (he inability to grow clinically relevant (he inability the inability of cells for infusion. An additional problem is that the inability administration of high doses of IL-2 necessary to maintain LAK activity

and CTL activity in vivo is associated with severe toxicity. Several techniques have been reported for improving the growth of cells for adoptive immunotherapy and for reducing the dosage requirement for systemic administration of IL-2. None of these attempts to increase activity provided a means to eliminate IL-2 from the protocol.

TIL cells activated with anti-CD3 mAb and expanded with moderate amounts of IL-2 (100 U/ml) have been successfully used in adoptive immunotherapy protocols using less toxic systemic doses of IL-2 [see, Goedegebuure, et al. (1995) J. Clin. Oncol. 13:1939, see, also, 10 Matsumura, et al. (1994) Cancer Research 54:2744]. In vivo administration of anti-CD3 mAb with low doses of IL-2 has also been suggested as an alternative adoptive immunotherapy strategy to lower the requirement for systemic IL-2 (see, Nakajima, et al. (1994) Proc. Natl. Acad. Sci. U.S.A. 91:78891. A method for expanding-CD4+-cells with 15 helper and cytolytic function using immobilized anti-CD3 mAb and IL-2 in rotary-tissue culture bags has also been described (see, Nakamura, et al. (1993) Br. J. Cancer 67:865]. Co-culture of anti-tumor effector cells activated with anti-CD3 mAb with lipopolysaccharide (LPS)-activated Bcells has also been suggested as an alternative method for growing cells 20 for adoptive immunotherapy [see, Okamoto, et.al. (1995) Cancer Immunol, Immunoth, 40:173]. These cells are all subsequently expanded with low doses of IL-2.

A combination of mAbs against CD3 and CD28 in the presence of lower dose IL-2 induces efficient expansion of TiL cells [see, Mulder, et al. (1995) Cancer Immunol Immunoth, 41:293]. Anti-tumor CTL generated by in vitro stimulation with synthetic peptides can grow as long as 4 months in culture with low dose IL-2 (30 u/ml), [see, Salgaller, et al. (1995) Cancer Research 55:4972). IL-7 has been shown to

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support the growth of CTL for prolonged periods in the absence of repeated stimulation [see, Lynch et al. (1994) J. Exp. Med. 179:31). Low concentrations of IL-2 have also been used to grow TIL cells in artificial capillary culture systems [see, Freedman, et al. (1994) J. Immunoth, Emphasis Tumor Immunot, 16(3):1981.

The need for exogenous IL-2 in expansion of immune cells has been obviated only by genetically modifying cells [see, e.g., U.S. Patent No. 5,470,730]. All the methods for growing genetically unmodified cells, however, require exogenous IL-2 to promote the differentiation and/or growth of cells for use in adoptive immunotherapy protocols. All methods require systemic administration of IL-2 to maintain activity of such cells.

Despite the showing of efficacy of adoptive immunotherapy in terminally-ill patients, the severe toxicity of the systematic dosages of fi15 2 required in adoptive immunotherapy protocols, the variability in the effector function of cell compositions derived from individual patients, as well as the difficulties in expanding clinically relevant numbers of effector cells has limited the use of adoptive immunotherapy. In particular, the need for exogenous IL-2 limits the cells used in adoptive immunotherapy to effector cells that can perform their functions over a limited period of time. In order to exploit the potential of this treatment method, there is a need to overcome the need for systemic IL-2 administration, and the difficulties in obtaining large quantities of cells. Thus, there is a need for improved adoptive immunotherapy methods.

Therefore, it is an object herein to provide such improved methods. In particular, it is an object herein to provide methods for expanding immune cells for use in adoptive immunotherapy protocols without the use of exogenous IL-2. It is also an object herein to provide methods to

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generate a large array of cell compositions, including compositions containing regulatory cells, for use in adoptive immunotherapy protocols. It is an object herein to provide means to produce compositions containing clinically relevant numbers of such cells. he availability of a an array of cell compositions permits the design of adoptive immunotherapy protocols for a wide variety of diseases and immune function alterations. Therefore, it is an object herein to provide methods for treating various disorders and altering immune function.

Compositions containing clinically relevant numbers of the immune cells are provided. The compositions contain regulatory immune cells, effector immune cells, or combinations thereof. In particular compositions containing, clinically relevant numbers of regulatory immune cells, especially Th1 and Th2 cells, for use in adoptive immunotherapy (herein referred to as autologous cell therapy (ACT)) are provided. Methods for generating the compositions containing the clinically relevant numbers of immune cells for use in adoptive immunotherapy are provided. The methods do not require use of IL-2. As a consequence, the expanded immune cells do not require LL-2 to retain activity of to remain viable.

Also provided are methods of treatment of disorders, including infectious diseases and, autoimmune diseases. In addition, methods of treatment for immunosuppression permitting organ or tissue transplantation and methods for enhancement of vaccination protocols are provided. The treatment methods use the compositions.

The compositions of regulatory cells provide a means to alter the immunoregulatory balance of a patient, either locally or sytemically, by changing the predominant regulatory cell population. Because many disease states occur with the loss of regulated balance of the immune system that is normally maintained by regulatory immune cells, the

availability of clinically-relevant numbers of regulatory immune cells provides a means to correct these imbalances. This ability offers great potential for treating a variety of diseases.

Methods for generating clinically relevant numbers of effector immune cells and of regulatory immune cells are provided. In particular, methods for generating substantially homogeneous populations of clinically relevant numbers of regulatory immune cells, including Th1 and Th2 cells, as well as Th1-like and Th2-like mononiclear cell populations are provided. Methods for generating compositions containing clinically relevant numbers of effector cells, such as CTLs, LAKS and TILS; that do not require exogenous IL-2 are provided.

Also provided are methods for producing clinically relevant quantities (Le., therapeutically effective numbers, typically greater than 10°, preferably greater than 10°) of autologous specific T cell types for treatment of disease states where a relative deficiency of such cells is observed. In particular, methods for producing clinically relevant numbers of autologous, ex vivo derived Th1 T-cells from patients with disease states where a Th2 cytokine profile predominates such as, but not limited to, infectious and allergic diseases; and autologous, ex vivo derived Th2 T-cells in Th1-dominant diseases, such as, but not limited to ,chronic inflammation and autoimmune diseases, for use in ACT protocols. The resulting cell compositions are provided and the use of the compositions in ACT protocols are provided.

Also provided are clinically relevant numbers of <u>ex vivo</u> derived

antigen-specific Th2 cells sensitized to a donor organ for use in ACT
protocols designed to provide specific immunosuppression for
transplantation procedures. Clinically relevant numbers of <u>ex vivo</u> derived
viral-specific Th1 cells for ACT protocols designed to provide protection

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from viral infection and thus serve as a viral vaccination strategy are also provided.

Methods of use of regulatory immune, cells in autologous cell therapy (ACT) protocols to treat and prevent human disease are provided. The ACT protocols designed to alter the immunoregulatory balance of a patient in order to treat diseases, where imbalances in regulatory cells exist. In particular, ACT protocols designed to alter the immunoregulatory, balance, of a patient in order to treat diseases where imbalances in regulatory cells exist are provided.

The methods involve collecting peripheral blood mononuclear cells from a patient and then expanding the cells by appropriate activation and then mitogenic stimulation with a cell surface specific proteins or proteins under conditions whereby clinically relevant numbers of the expanded cell type are produced (typically 10°, preferably 10°, more preferably 10°), or more depending upon the cell type and ultimate application). If the collected cells are not differentiated in vivo or require further differentiation, then following collection and prior to expansion, the method includes activating and causing differentiation of the cells ax vivo under conditions whereby at least some of the cells differentiate into regulatory or effector cells or other cell types. The resulting cells are then reinfused into the donor to effect treatment. The desired cells may be purified prior to reinfusion to provided a more homogeneous population.

Where required, differentiation of mononuclear cells is effected by activating the cells with a mitogen in the presence of the appropriate array of cytokines. This activation can be achieved by use of agents, such as cytokines or mitogens or other growth promoting agents under environmental conditions conducive to development of a particular phenotype. For example, if the cells are activated in the presence of IFN-y, Th1 cell differentiation will be produced. If they are activated in

the presence of IL-4, then Th2 cell differentiation will be produced. Such activating agents include monoclonal antibodies for polyclonal activation, and natural or synthetic antigens for specific activation presented in the context of MHC molecules.

Expansion is effected by growing the cells under conditions in which high cell densities can be achieved, whereby endogenous cytokines will be retained in the vicinity of the growing cell population, and in the presence of one or more mitogenic monoclonal antibodies or other cell surface specific protein, other than IL-2 or other such cytokine that will require co-infusion. Such conditions are preferably achieved by growing the cells in a hollow fiber (HF) bioreactor.

Methods for treating various disorders using the resulting cells are also provided. In effecting these methods, cells of a type that are found to be deficient or in low relative amounts are infused into a patient. For example, infectious diseases or tumors may be treated by collecting peripheral blood mononuclear cells from a patient; expanding the cells under conditions whereby a composition containing a therapeutically effective number of cells is produced; and infusing the resulting composition of cells into the patient. In preferred embodiments, the cells are specific for unique antigens in the vicinity of the site where an effect is desired or are specific for a pathogen or tumor being treated. In other preferred embodiments, effector cells, such as cytotoxic CD8.³

T. lymphocytes (CTLs) that are specific for the pathogen or tumor are infused or co-infused with regulatory cells.

In addition, methods for specific immunosuppression for transplantation procedures are provided. These methods involve administration of clinically relevant numbers of <u>ex vivo</u> derived antigenspecific Th2 cells sensitized to a donor organ. In preferred embodiments the cells are specific for alloantigens or an antigen unique to the organ or tissue being transplanted.

Also provided are vaccination methods and compositions for use as vaccines. In particular the vaccines are formulated from clinically relevant numbers of ax vivo-derived viral-specific Th1-cells or Jh2 cells (or Th1-like or Th2-like populations of cells) that upon infusion provide protection from viral infection and thus serve as a viral vaccination strategy.

infusing autologous, ax vivo derived and expanded regulatory immunes of the provided. This method includes the steps of collecting peripheral blood mononiclear cells from a patient, activating the cells ax vivo under conditions whereby at least some, even one, of the cells differentiate into the desired regulatory cells, expanding the regulatory cells, and infusing the expanded regulatory cells into the donor to affect the immunoregulatory balance. In particular, the infusion is not accompanied

by co-infusion of a cytokine, such as IL-2.

The method above is useful for therapeutic treatment of disorders characterized by imbalances in regulatory immune cells. Specifically, the methods provided herein can be used to develop treatments, for chronic inflammation in disorders such as, but not limited to; multiple sclerosis; rheumatoid arthritis, Crohn's Disease, autoimmune thyroid disease and inflammatory bowel disease; chronic infectious diseases such as infections with human immunedeficiency virus, herpes simplex virus, cytomegalovirus and hepatovirus; allergic and other hypersensitivity disorders such as asthma; and provides a method for specific immunosuppression in organ and tissue transplant procedures and a method to provide immunoprotection in vaccination.

In preferred embodiments, the regulatory immune cells are either Th1, Th2 or Th3 cells with a CD4+ or CD8+ phenotype. The cells will

preferably have a "memory" phenotype (i.e., CD45RO*, L-selectin), which permit the cells to traffic to sites of inflammation. These cells are preferably made to exert their regulatory function at a localized area of the body by selectively expanding cells specific for an unique antigen present at the site the regulatory effect of the cells is desired. For example, in the treatment of rheumatoid arthritis, regulatory cells specific for type II collagen, which is present only in joint tissue, are preferred. In the treatment of diabetes for preventing rejection of transplanted islet cells, regulatory cells specific for insulin are preferred.

10 In other embodiments, the cells are effector cells that have been expanded up to clinically relevant (<u>i.e.</u>, therapeutically effective) numbers without the use of IL-2 to promote expansion.

Also provided is a method for expanding immune cells without the use of exogenous IL-2. The expansion of immune cells is preferably caused by the inclusion of one or more mitogenic mAb in the culture medium. The immune cells are preferably expanded under conditions in which they grow to high density. Such high density can be achieved by growing the cells in hollow fiber bioreactors with the molecular weight cut-offs of the fibers that retain endogenously produced cytokines. Such molecular weigh cut-off is preferably less than 14,000 daltons, more preferably 6000 daltons.

Also provided are methods for producing clinically relevant populations of virally purged CD4* cells obtained from HIV* patients.

The resulting virally purged CD4* cells are then reinfused into the donor patient in order to effect treatment of HIV. The cells may also be co-infused with anti-HIV effector cells.

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DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

A. Definitions

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as is commonly understood by one of skill in the art to which this invention belongs. All patents and publications referred to herein are, unless noted otherwise, incorporated by reference in their entirety.

As used herein, adoptive immunotherapy or cellular adoptive immunotherapy refers to a method of treatment involving administration of immunologically active cells. The cells used in the treatment are generally obtained by venipuncture or leukopheresis either from the individual to be treated (autologous treatment) or from another individual (allogeneic). For purposes herein, autologous treatment is herein referred to as autologous cell therapy (ACT).

As used herein, autologous cell therapy [ACT] is a therapeutic method in which cells of the immune system are removed from an individual, cultured and/or manipulated <u>ex vivo</u> or <u>in vitro</u>, and introduced into the same individual as part of a therapeutic treatment.

As used herein, activating proteins are molecules that when contacted with a T-cell population cause the cells to proliferate. T-cells generally require two signals to proliferate. Activating proteins thus encompasses the combination of proteins that provide the requisite signals, which include an initial priming signal and a second costimulatory signal. The first signal requires a single agent, such as anti-CD3 mAb, anti-CD2 mAb, enti-TCR mAb, PHA, PMA, and other such signals. The second signal requires one or more agents, such as anti-CD28, anti-CD40L, cytokines and other such signals. Thus activating proteins include combinations of molecules including, but are not limited to: cell surface protein specific monoclonal antibodies, fusion proteins

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containing ligands for a cell surface protein, ligands for such cell surface proteins, or any molecule that specifically interacts with a cell surface receptor on a mononuclear cell and indirectly or directly causes that cell to proliferate. For purposes herein, when expanding effector cells, the 5 activating proteins are selected from among those that are not needed to substantially maintain cell viability and function after expansion. Thus for example, IL-2 is not an activating protein for purposes berein for the effector cell expansion. As noted, the methods herein provide a means to produce cells, particularly effector, that do not require IL-2, and thus, in preferred embodiments, IL-2 will not be used as an activating agent,

As used herein, a mitogenic monoclonal antibody is an activating protein that is an antibody that when contacted with a cell directly or indirectly provides one of the two requisite signals for T-cell mitogenesis: Generally such antibodies will specifically bind to a cell surface receptor thereby inducing signal transduction that leads to cell proliferation. Suitable mitogenic antibodies may be identified empirically by testing an selected antibodies singly or in combination for the ability to increase numbers of a specific effector cell. Suitable mitogenic antibodies or combinations thereof will increase the number of cells in a selected time 20 period, typically 1 to 10 days, by at least about 50%, preferably about 100% and more preferably 150-200% or more, compared to the numbers of cells in the absence of the antibody.

As used herein, a growth promoting substance is a substance, that may be soluble or insoluble, that in some manner participates in or the second induces cells to differentiate, activate, grow and/or divide. Growth promoting substances include mitogens and cytokines. Examples of growth promoting substances include the fibroblast growth factors, at the 1 osteogenin, which has been purified from demineralized bone [see, e.g., Luyten, et al. (1989) J. Biol. Chem. 264:13377)), epidermal growth

factor, the products of oncogenes, the interleukins, colony stimulating factors, and any other of such factors that are known to those of skill in the art. Recombinantly-produced growth promoting substances, such as recombinantly-produced interleukins, are suitable for use in the methods herein. Means to clone DNA encoding such proteins and the means to

nergin. Means to clone UNA engoding such proteins and the means to produce biologically active proteins from such cloned DNA are within the skill in the art. For example, interleukins 1 through 6 and others have been cloned. Various growth promoting substances and combinations thereof may be used to expand desired subpopulations of lymphoid cells.

divide and in particular, a mitogen is a substance that induces cells to divide and in particular, as used herein, are; substances that stimulate as lymphocyte population in an antigen-independent manner to proliferate and differentiate, into effector cells or regulatory cells. Examples of such substances include lectins and lipopolysaccharides.

As used herein, a cytokine is a factor; such as lymphokine or monokine, that is produced by cells that affect the same or other cells!

As used herein, a lymphokine is a substance that is produced and secreted by activated T lymphocytes and that affects the same or other cell types. Tumor, necrosis factor, the interleukins and the interferons are examples of lymphokines. A monokine is a substance that is secreted by monocytes or macrophages that affects the same or other cells.

As used herein, a regulatory immune cell is any mononuclear cellwith a defined cytokine production profile and in which such cytokine
profile does not directly mediate an effector function. A regulatory

immune cell is a mononuclear cell that has the ability to control or direct
an immune response, but does not act as an effector cell in the response.

Regulatory immune cells exert their regulatory function by virtue of their cytokines
production profile. For example, regulatory immune cells that produce the

2 and IFN-y, but do not produce IL-4 are termed "Th1" cells. Regulatory immune cells that produce IL-4 and IL-10, but do not produce IFN-y are termed "Th2", cells. Regulatory immune cells that produce IFN-y are termed "Th2", cells. Regulatory immune cells that produce TGF-p, IL-10 and IFN-y, but do not produce IL-2 or IL-4 are termed "Th3" cells. Cells that produce Th1, Th2 and Th3 cytokine profiles occur in CD4+ and CD8+ cell populations. Cells that produce "IL-2, IL-4' and IFN-y are thought to be precursors of Th1" and Th2 cells and are designated "Th0" cells. Populations of cells that produce a majority of Th1" cytokines are designated Th2-like"; those producing a majority of Th3 cytokines are designated Th3-like"; those producing a majority of Th3 cytokines are designated "Th3-like". Thus, each composition, although containing a heterogeneous population of cells, will have the properties that are substantially similar, with respect to cytokine, to the particular Th subset.

It is understood that this list of T- cells is exemplary only, and any other definable population, array or subtype of T-cells that can be expanded by the methods herein to clinically relevant humbers are intended herein.

As used herein, a composition containing a clinically relevant number or population of immune cells is a composition that contains at least 109, preferably greater than 109, more preferably at least 1010 cells, and most preferably more than 1010 cells, in which the majority of the cells have a defined regulatory or effector function; such as Th1 cells or Th2 cells or effector cells, such as LAK, TIL and CTL cells. The preferred number of cells will depend upon the ultimate use for which the composition is intended as will the type of cell. For example, if Th1 cells that are specific for a particular antigen are desired, then the population will contain greater than 50%, preferably greater than 70%, more preferably greater than 70%, more

such cells. If the population results from polyclonal expansion, the homogeneous cells will be those that are a particular type or subtype. For uses provided herein, the cells are preferably in a volume of a liter or less, more preferably 500 mls or less, even more preferably 250 mls or less and most preferably about 100 mls or less.

As used herein, predominant means greater than about 50%.

As used herein, a combination refers to two component, items, such as compositions or mixtures, that are intended for use either together or sequentially. The combination may be provided as a mixture of the components or as separate components packaged or provided together, such as in a kit.

As used herein, effector cells are mononuclear cells that have the ability to directly eliminate pathogens or tumor cells. Such cells include, but are not limited to, LAK cells, MAK cells and other mononuclear phagocytes... Tiles, CTLs and antibody-producing B cells and other such cells.

As used herein, immune balance refers to the normal ratios, and absolute numbers, of various immune cells that are associated with a disease free state. Restoration of immune balance refers to restoration to a condition in which treatment of the disease or disorder is effected whereby the ratios of regulatory immune cell types and numbers thereof are within normal range or close enough, thereto so that symptoms of the treated disease or disorder are ameliorated. The amount of cells to administer can be determined empirically, or, preferably, by administering aliquots of cells to a patient until the symptoms of the disease or disorder are reduced or eliminated. Generally a first dosage will be at least 10° 10° cells. In addition, the dosage will vary depending upon treatment as sought.

As used herein, therapeutically effective refers to an amount of cells that is sufficient to ameliorate, or in some manner reduce the symptoms associated with a disease. When used with reference to a method, the method is sufficiently effective to ameliorate, or in some manner reduce the symptoms associated with a disease.

As used herein, mononuclear or lymphoid cells (the terms are used interchangeably) include lymphocytes, macrophages, and monocytes that are derived from any tissue in which such cells are present. In general lymphoid cells are removed from an individual who is to be treated. The 10 lymphoid cells may be derived from a tumor, peripheral blood, or other tissues, such as the lymph nodes and spleen that contain or produce lymphoid cells

As used herein, therapeutically useful subpopulations of in vitro or ex vivo expanded mononuclear or lymphoid cells are cells that are 15 expanded upon exposure of the cells to a growth promoting substances. such as lymphokines, when the lymphoid cells are cultured ex vivo. therapeutically useful subpopulations are regulatory cells or effector cells and contain clinically relevant numbers of cells, typically at least about 109 or more cells, which are preferably in a clinically useful volume (i.e., for infusion) that is one liter or less.

As used herein, a therapeutically effective number or clinicallyrelevant number ex vivo expanded cells is the number of such cells that is at least sufficient to achieve a desired therapeutic effect. when such cells are used in a particular method of ACT. Typically such number is at 25 least 109, and more preferably 1010 or more. The precise number will depend upon the cell type and also the intended target or result.

As used herein, a hollow fiber bioreactor or hollow fiber bioreactor cartridge contains an outer shell casing that is suitable for the growth of mammalian cells, a plurality of semi-permeable hollow fibers encased within the shell that are suitable for the growth of mammalian cells on or near them, and the ECS, which contains the cells and the ECS cell and supernatant. The interior of the hollow fibers is called the lumen and the area between the outside of the capillaries to the inside of the outer housing is called the extracapillary space (ECS).

Tissue culture medium perfuses through the fiber lumens and is also included within the shell surrounding said fibers. The tissue culture medium, which may differ in these two compartments contains diffusible components that are capable of sustaining and permitting proliferation of immune cells. The medium is provided in a reservoir from which it is pumped through the fibers. The flow rate can be controlled varied by the varying the applied pressure. The ECS or perfusing medium may additionally contain an effective amount of at least one growth promoting or suppressing substance that specifically promotes the expansion or suppression of at least one subpopulation of the immune cells, such as: Til. cells or regulatory cells, in which the effective amount is an amount 20 sufficient to effect said specific expansion.

As used herein, a hollow cell fiber culture system includes of a hollow fiber bioreactor as well as pumping means for perfusing medium through said system, reservoir means for providing and collecting medium, and other components, including electronic controlling, recording or sensing devices. A hollow fiber bioreactor is a cartridge that contains of a multitude; of semi-permeable tube-shaped fibers encased in a hollow shell. The terms hollow fiber reactor and hollow fiber bioreactor are used interchangeably. A preferred device for methods is that described in copending, allowed, U.S. application Serial No. 08/506, 173.

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As used herein, ECS refers to the extra-capillary space cell supernatant. It is the medium in which the cells in the ECS are growing. It contains secreted cellular products, diffusible nutrients and any growth promoting or suppressing substances, such as lymphokines and cytokines, produced by the cultured immune cells or added to the ECS or tissue culture medium. The particular components included in the ECS is a function not only of what is inoculated therein, but also of the characteristics of the selected hollow fiber.

As used herein, tissue culture medium includes any culture medium that is suitable for the growth of mammalian cells ex vivo. Examples of such medium include, but are not limited to AIM-V, RPMI 1640, and Iscove's medium (GIBCO, Grand Island, N.Y.). The medium may be supplemented with additional ingredients including serum, serum proteins, growth suppressing, and growth promoting substances, such mitogenic monoclonal antibodies and selective agents for selecting genetically engineered or modified cells.

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As used herein, treatment means any manner in which the ""
symptoms of a condition, disorder or disease are ameliorated or otherwise
beneficially altered. Treatment also encompasses any pharmaceutical use
of the compositions herein.

As used herein, a vaccine is a composition that provides protection against a viral infection, cancer or other disorder or treatment for a viral infection, cancer or other disorder. Protection against a viral infection, cancer or other disorder will either completely prevent infection or the tumor or other disorder or will reduce the severity or duration of infection, tumor or other disorder if subsequently infected or afflicted with the disorder. Treatment will cause an amelioration in one or more symptoms or a decrease in severity or duration.

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As used herein, amelioration of the symptoms of a particular disorder by administration of a particular composition refers to any lessening, whether permanent or temporary, lasting or transient that can be attributed to or associated with administration of the composition.

As used herein, substantially pure means sufficiently homogeneous to appear free of readily detectable impurities as determined by standard methods of analysis, such as flow cytometry; used by those of skillin the art to assess such purity, or sufficiently pure such that further purification would not detectably alter the physical and chemical properties, such as biological activities, of the substance. Methods for purification of the immune cells to produce substantially pure populations are known to those of skill in the art. A substantially pure cell population, may, however, be a mixture of subtypes; purity refers to the activity profile of the population. In such instances, further purification might increase the specific activity of the cell population.

As used herein, biological activity refers to the in vivo activities of immune cells or physiological responses that result upon in vivo administration of a cell, composition or other mixture... Biological activity, thus, encompasses therapeutic effects and pharmaceutical activity of such cells, compositions and mixtures.

Although any similar or equivalent methods and materials can be employed in the practice and/or tests of the methods and cells provided herein, preferred embodiments are now described.

B. Effector and regulatory immune cells

Encounter of a host with antigen can result in either cell-mediated or humoral classes of immune response. Regulatory immune cells control the nature of an immune response to pathogens [see, Mosmann, et al. (1986) J. Immunol. 136:2348; Cherwinski, et al. (1987) J. Exp. Med. 166:1229; and Del Prete, et al. (1991) J. Clin. Invest, 88:346]. The

different types of responses are attributable to the heterogeneity of CD4* Colls. CD4* cells can be sub-divided according to their cytokine expression profiles. These cells are derived from a common precursor, Th0, which can produce Th1; Th2 and Th3 cytokines [see, Firestein, et al. (1989) J. Immunol. 143:518]. As noted above, Th1 clones produce IL-2, INF-y, lymphotoxin and other factors responsible for promoting delayed-type hypersensitivity reactions characteristic of cell-mediated immunity. These cells do not express IL-4 or IL-5. Th1 cells promote cell-mediated inflammatory reactions, support macropriage activation, immunoglobulin (Ig) isotype switching to IgG2e and activate cytotoxic function.

Th2 clones produce cytokines, such as IL-4, II-5, IL-6, IL-10 and IL-13, and thus direct humoral immune responses, and also promote allergic type responses. Th2 cells do not express IL-2 and IFN-y. Th2 cells provide help for B-cell activation, for switching to the IgG1 and IgE isotypes and for antibody production [see, e.g., Mosmann et al. (1989) Annu. Rev. Immunol. Z:145]. Th3 cell produce IL-4; IL-10 and TGF-8. The cytokines produced by Th1 and Th2 cells are mutually inhibitory. Th1 cytokines inhibit the proliferation of Th2 cells and Th2 cytokines inhibit Th1 cytokine synthesis [see, e.g., Florentino, et al. (1989) Med. 170:2081 (1989). This cross regulation results in a polarized Th1 or Th2 immune response to pathogens that can result in host resistance or susceptibility to infection.

Development of the appropriate regulatory immune cell response during infection is important because certain pathogens are most effectively controlled by either a predominantly Th1 or Th2 type immune response [see, e.g.], Sher, et al. (1989) Ann. Rev. Immunol. 46:111; Scott, et al. (1991) Immunol. Today 12:346; Sher, et al. (1992) Immunol. Rev., 127:183; and Urban, et al. (1992) Immunol. Rev.

127:205]. For example, a correlation has been found between the predominant regulatory immune response and disease susceptibility in leprosy [see, e.g., Yamamure, et al. (1991) Science 254:277] AIDS [see, e.g., Clerici, et al. (1993) Immunol. Today 14:107], toxoplasma [see,

- Sher, et al. (1989) Ann. Rev. Immunol. 48:111], Hashimoto's thyroiditis [see, e.g., Del Prete, et al. (1989) Autoimmunity 4:267], Grave's disease [see, e.g., Turney, et al. (1987), Eur. J. Immunol. 47:1807], transplantation [see, e.g., Benyenuto, et al. (1991) J. Pathol. 11:887], type 1 diabetes [see, e.g., Foulig, et al. (1991) J. Pathol. 11:55:97], multiple sclerosis [see, e.g., Benyenuto, et al. (1991) Clin Exp.
- 10 165:97), multiple scierosis [see, e.g., Benvenuto, et al. (1991) Clin. Exp. Immunol. 84:97), and rheumatoid arthritis [see, e.g., Quayle, et al. (1993) Scand. J. Immunol 38:75].
- A Th1 response in mice to protozoan, viral and fungal infection is associated with resistance, while a Th2 response is associated with the saccitated with the saccit
- Subsets of CD8* T-cells also are known to secrete a Th1- or Th2cytokine pattern. Exposure of CD8* cells to IFN-y and IL-2 direct
 differentiation into Th1 cells; whereas, IL-4 induces differentiation into
 Th2 cells. Th1 CD8* cells are thought to be important effectors in the
 immune response to viruses, while Th2 CD8* cells have an immuno-

Immunology Today 17:138-146].

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suppressive function. Other regulatory cells can be characterized by methods similar to those used to characterize the above described cells.

By virtue of the cross regulation and the immune imbalances observed in disease states, as described herein, regulatory cells should be therapeutic for the treatment of a variety of diseases. Such use has been demonstrated to some extent in animal models, but has not been possible to achieve in humans. For example, administration of native T-cells and The antigen-specific clones for Actinobacilitis actinomycetemicomitans, in combination did amidiorate periodontal disease in nude rats (see, 1994). Antigen-specific Thi cellschops have been shown for intract analyst intertific.

specific Th1 cell clones have been shown to protect against infection with the protozoan Leishmania major, genital infection with Chiamydia trachomatis and murine candiclasis (see, Powile, et al. (1994) 1. Exp. Med. 179:589; Igletseine; et al. (1993) at al. Regional immunity 5:317; and Romani (1991) Interiment, 59:4647). In addition, Th2 cell clones have been shown to prevent autoimmune everetinitis (see Saoudi, et al. (1993) Eur. 1. Immunol; 23:3096). An antigen specific Th2 cell clone has been shown to suppress an animal model of multiple scienosis (see, Chen. et al. (1994) Science 265:1237). Donor-specific Th2 cells can reduce lethal graft vs. host disease in transplantation (see, Fowler, et al. (1994) Adv. Bone Marrow Purg. Process, Fourth Int. Sympos., Wiley-Liss, Inc., p. 533. Purified T-cells with enhanced Th2 activity have also been shown to prevent insulin-dependent diabetes-like disease in animals. See, Fowell et al. (1993) J. Exp. Med. 177:627.

While Th2 clones have been used in adoptive transfer studies in animals, regulatory cells, including Th1 and Th2 cells, have not been used in ACT protocols in humans. Such protocols are limited by the inability to differentiate and produce therapeutically effective quantities of such regulatory cells. The methods herein, however, provide a means to

produce such clinically relevant quantities of cells, and, thereby provide a means to ameliorate disorders, provide vaccines, and suppress tissue or organ rejection. The methods herein also provide a means to produce clinically relevant quantities of relulatory and effector cells in the absence of IL-2.

Also provided herein, are methods for growing cells that are the appendically useful, for treatment of HIV infection, including treatment of A.I.D.S. by enchancing or restoring the immune system [see, e.g., see Examples 3 and 4].

10 C. Methods for production of regulatory cells

A method for obtaining regulatory cells for use in ACT protocols is provided herein. A method for obtaining effector cells for use in ACT protocols without the need for exogenous agents, such as IL-2, that sustain the viability of such cells is also provided. The method includes some or all of the following steps: (1) collecting mononuclear cells from a patient; (2) treating the cells <u>ex vivo</u> with that agents that cause some or all of the cells to the differentiate into desired T cell subtypes; (3) purifying the resulting cells; and (4) expanding these cells by contacting them with a mitogenic agent that specifically interacts with a cell surface receptor. Such agents are herein preferably mitogenic monoclonal antibodies. The expanded cells may be further purified to select for the desired subtype.

1. Collecting mononuclear cells

Mononuclear cells (i.e., lymphocytes and monocytes) can be obtained from a variety of sources, including, but not limited to peripheral blood, lymphoid tissue, biopsy tissue or from body cavity lavage procedures. Preferably, the cells are obtained by simple venipuncture (50-500 ml). When larger numbers of cells are required, they may be obtained by a lymphapheresis procedure. The mononuclear

cells can be purified from the blood using Ficoll-Hypaque density gradient centrifugation or any other suitable method.

a. Ex Vivo Differentiation

- Many studies have indicated that different antigens can cause a selective induction of distinct immunoregulatory cell subsets, causing the development of either a humoral or cell-mediated immune response. Furthermore, many disease states are the result of the predominance of the certain cell types. Recent advances in the understanding of the mechanisms regulating the differentiation of T-cell subsets allows the generation of selected subsets ex vivo.
- presenting cell and the MMC haplotype of an individual can affect the differentiation of specific types of regulatory immune cells. Various cytokines are also able to affect the type of regulatory response that develops in a person. For example, it is known that the presence of IL-4 during initial T-cell activation gives rise to Th2-like cells (see, Hsieh, et al. (1992) Proc. Natl. Acad. Sci. U.S.A. 89:6065 and Paliard, et al. (1988) et al. J. Immunol. 141:849]. Conversely, activation of cells in the presence of IL-12 or interferon-gamma leads to the formation of Th1-like cells (see, Sedar, et al. (1993) Proc. Natl. Acad. Sci. U.S.A. 90:10188).
 - Accordingly, in a preferred embodiment, the mononuclear cells collected in the first step of the present process are next activated in the presence of IL-12, interferon-gamma or IL-4 to cause the development of Th1 or Th2 cells, respectively. To enhance the differentiation of
- 25 regulatory cells, antibodies to IL-12 and/or interferon gamma can be used to promote Th2 responses, while antibodies to IL-4 can be used to promote the differentiation of Th1 cells. Antibodies or other proteins specific for the IL-12, interferon-gamma or IL-4 receptor on T-cells could also be used to provide a signal in place of the lymphokines. The cells

can be activated either non-specifically with chemical agents such as PHA and PMA or with monoclonal antibodies such as anti-CD3, or anti-CD2; ... Preferably, they are activated specifically with natural or man-made protein antigens added to the medium, processed and presented by APC to T-cells. It may be necessary in some cases to vaccinate the patient prior to blood collection in order to increase the starting number of antigen-specific cells. Another strategy is to oral tolerize patients prior to blood collection. In cases where the cells generated are specific for a known antigen, the antigen may also be used after the cell reinfusion as a booster to increase the desired regulatory cells in vivo. Additional and increase the desired regulatory cells in vivo. strategies for effecting Th1 cell differentiation is to activate cells in the presence of qB7.2 mAb or TGF-β. Th2 differentiation also can be promoted by activating cells in the presence of one or more of agents. such as, one or more of the following: aB7.1 mAb, low antigen doses 15 and CTLA4/lg fusion protein (CTLA4 is a ligand for CD28); CD28 is expressed on Tacells and antigen presenting cells above 10.10 to a 10.00 m. The type of regulatory cells generated should be determined from animal models of the disease. It is known that not all regulatory cells within a classification are alike. For example, some Th2 cells secrete high 20 levels of IL-4 and low levels of IL-10, while others have increased levels of IL-5. Other regulatory cells produce IL-10 and interferon-gamma. Regulatory cells termed "Th3" cells secrete TGF-β and are deemed preferential for treatment of multiple sclerosis. 1000

b. Regulatory Cell Isolation

Most techniques for isolation of immune cell subsets are based on the reactivity of mAb against T-cell surface antigens. Positive selection can be achieved by fluorescent-activated cell sorting [see, Reinherz, et al. (1979) Proc. Natl. Acad. Sci. U.S.A. 76:4061]. Various panning techniques where specific mAb are bound to plastic plates to capture the

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desired T-cell subsets can also be used. See, Lum, et al. (1982) Cell Immunol. 72:122.

Panning techniques can be used for negative selection as well, depleting unwanted subsets with specific mAb [see, s.g., Engleman, st al. (1981) J. Immunol. 127:2124]. The use of magnetic polymer beads coated with mAb is a preferred method to isolate highly purified, functionally intact lymphoid cell populations by positive and negative selection [see, sig., Lea, et al. (1985) Scand J. Immunol. 22:207, Lea, et al. (1986) Scand J. Immunol. 23:509) and Gaudernack, et al. (1986) J. Immunol. Methods 90:179).

regulatory immune cell subsets, efforts must be made to enhance the desired population by purifying on the basis of certain cell surface proteins. For example, CD30 positive (see! Manetti, et al. (1994) J. Exp. Med. 180:2407); CD27 negative (see, Elson, et al. (1994) Int. Immunol. 6:1003] and CD7 negative (see, Autran, et al. (1995) J. Immunol. 154:1408] cell populations have been shown to have the majority of Th2 cells. Also, repeatedly contacting the cells with anti-CD28 mAb is another method for enhancing Th2 cells.

Another strategy for purification of regulatory cells is to expand the cells in the presence of agents known to inhibit the growth of the unwanted subset(s) of cell. Such agents include dexamethasone, colchicine, CTLA4/lg fusion protein and progesterone, which inhibit Th2 cell growth. TGF-g inhibits Th1 cell growth.

c. Regulatory Cell Expansion

Methods for expanding purified T-cells to clinically relevant numbers ex vivo without the use of exogenous |L-2 are provided herein.

Although |L-2 could be used in the present methods, it is preferably to grow cells without the addition of this cytokine. Cells exposed to ||-2 ex

viso may become dependent on the presence of IL-2 to maintain their visibility and function, requiring the systemic infusion of IL-2 with the cells to the patient. Because the systemic infusion of IL-2 is known to be extremely toxic to patients it is best to avoid the necessity for this blood cytokine.

In order for T-cells to proliferate, they require two separate signals.

The first signal is generally delivered through the CD3/TCR antigen complex on the surface of the cells. The second is generally provided through the IL-2 receptor. In order to bypass the IL-2 receptor. In order to bypass the IL-2 receptor to combinations of mAb are used. Preferably, the mAb are in the soluble in the so

phase or immobilized on plastic or magnetic beads; inforder to simplify the cell harvesting procedure; at the cell harvesting procedure; at the cell harvesting procedure; at the cell harvesting of the cell harvesting procedure; at the cell harvesting of the cell harvesting o

To, provide the first signal, it is preferable to activate cells with

mAb, to, the CD3/TCR complex, but other suitable signals, such as; but
not limited to, antigens, super antigens, polyclonal activators; anti-CD2
and anti-TCR antibodies, may be used. Other suitable agents can be
empirically identified.; Immobilized or cross-linked anti-CD3 mAb, such as

OKT3, or 64.1, can activate T-cells in a polyclonal manner (see, Tax, et al.

1983) Nature 304:445). Other polyclonal activators; however, such as
phorbol myristate acetate can also be used [see, e.g., Hansén] et al.

(1980) Immunogenetics 10:247).

Monovalent anti-CD3; mAb in the soluble phase can also be used to activate T-cells (see, Tamura, et al. (1992) J. Immunol. 148:2370).

Stimulation of CD4 + cells with monovalent anti-CD3 mAb in the soluble form is preferable for expansion of Th2 cells, but not Th1 cells (see, and deJong, et al. (1992) J. Immunol. 149:2795]. Soluble heteroconjugates of anti-CD3 and anti-T-cell surface antigen mAb can preferentially activate a particular T-cell subset (see, Ledbetter, et al. (1988) Eur. S.

Immunol. 18:525]. Anti-CD2 mAb can also activate T-cells [see, Huet, et al. (1986) J. Immunol. 137:1420]. Anti-MHC class II mAb can have a synergistic effect with anti-CD3 in inducing T-cell proliferation [see. Spertini, et al. (1992) J. Immunol. 149:65]. Anti-CD44 mAb can activate 5 T-cells in a fashion similar to anti-CD3 mAb. See, Galandrini, et al.

(1993) J. Immunol. 150:42251.68 (1993) 4 (1993) 4 (1993)

For purposes herein, monoclonal antibodies to anti-CD3 are preferred. Anti-CD3 is used because CD3 is adjacent to the T-cell receptor. Triggering of CD3, such as by monoclonal antibody interaction. 10 causes concomitant T cell activation. (a) The second of the second activation.

version of markii). Second signal mark at the real total larger to a set the To then cause proliferation of such activated T cells, a second signal is required. A variety of mAb singly or in combination can provide the second signal for T-cell proliferation. Anti-IL-4R mAb (specific for the interleukin-4 receptor molecule) can enhance the proliferation of the Th2 cells [see, Lindquist, et al. (1993) J. Immunol, 150:3941. Immobilized ligands or mAb against CD4, CD8, CD11a (LFA-1), CD49 (VLA) CD45RO, CD44 and CD28 can also be used to enhance T-cell proliferation [see, Manger, et al. (1985) J. Immunol. 135:3669; Hara, et 20 al. (1985) J. Exp. Med. 161:1513; Shimizu, et al. (1990) J. Immunol. 145:59; and Springer, (1990) Nature 346:425). Cell surface proteins that are ligans to B-cells are preferred targets for Th2 cell proliferation.

Anti-CD28 mAb in combination with anti-CD3 or anti-CD2 induces a long lasting T-cell proliferative response [see, Pierres, et al. (1988) Eur. J. Immunol. 18:685]. Anti-CD28 mAb in combination with anti-CD5 mAb results in an enhanced proliferative response that can be sustained for weeks [see, Ledbetter, et al. (1985) J. Immunol. 135:23311. Anti-CD5 mAb alone can also provide a second signal for T-cell proliferation

while macrophage ligands are preferred for Th1 cell proliferation.

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[see, Vandenberghe et al. (1991) Eur. J. Immunol. 21:251]. Other mAb known to support T-cell proliferation include anti-CD45 and CD27 [see, Ledbetter, et al. (1985) J. Immunol. 135:1819 and Van Lier, et al. (1987) J. Immunol. 139:1589).

To determine the combination of mAbs or proteins that optimally, induce sustained regulatory cell proliferation, a screening procedure using combinations of these mAbs or proteins is used. The cells are incubated with various combinations of these substances and screened for growth by analysis of 3H-thymidine, incorporation or equivalent methods. The group demonstrating the best growth characteristics is selected for use in the medium.

(iii) Expansion

In order to expand purified T-cells to clinically relevant numbers of up to 100 billion (101), the cells should be grown to high density. This can be achieved using any suitable means, including, but not limited to: stirred tank fermentors, airlift fermentors, roller bottles, culture bags, and other bioreactor devices. Hollow fiber bioreactors are presently preferred. Hollow fiber bioreactors permit cells to be cultured to the required high densities in a minimal volume. This reduces the amount of monoclonal antibodies, serum and medium required in the production process. In addition, selection of fibers with molecular weight cut-offs of 6000 addition, selection of fibers with molecular weight cut-offs of 6000 additions, will allow continuous feeding and waste product removal while retaining cell derived cytokines in the culture space. These cytokines, such as IL-2 and IL-4, promote and sustain cell viability and proliferation.

T-cells, like most mammalian cells, will grow to a maximum density of 1 x 10⁶ cells/ml in tissue culture. Thus, a total of 100 liters of culture medium would be required to support 100 billion cells. In addition, there 100 liters of medium would have to be replenished regularly to maintain a proper nutrient/waste product balance necessary to keep the cells viable.

A method would also be required to keep the 100 liters of medium saturated with oxygen.

Hollow fiber technology for cell culture is well known [see, e.g. U.S. Patent Nos. 4,220,725, 4,206,015, 4,200,689, 3,883,393, and 5 3.821,087; see, also, U.S. Patent No. 4,391,912; U.S. Patent No. 4,546,083; U.S. Patent No. 4,301,249; U.S. Patent No. 4,973,558, U.S. Patent No. 4,999,298: and U.S. Patent No. 4,629,6861 and is used to achieve issue-like cell densities in culture (i.e., densities of greater than about 108 cells/ml]. The original hollow fiber bioreactor contains a housing with a plurality of artificial capillary hollow fiber membranes. The capillaries extend between an inflow opening at one end of the device and an outflow opening at the other. The capillaries have selectively permeable walls though which dissolved medium components can diffuse. The lumen and ECS are separated by potting material at the inflow and outflow openings. The housing also contains ports for access 15 to the ECS enabling cells to be inoculated into the ECS [see, e.g., U.S. Patent Nos. 3,821,087; 3,883,393 and 4,220,725, 4,206,015. 4,200,689, 3,883,393, and 3,821,087; see, also Knazek, et al. (1972) Science 178:651.

Hollow fiber technology permits cells to grow to densities 100-fold greater than cell densities [1 x 108 cells/ml or greater] observed in conventional cell culture. Thus, only one liter of culture volume is required to generate 100 billion cells. The reduced cell volume would also decrease the amount of human serum and soluble mAb required in 25 the expansion process. In addition, high cell densities provide environments that are a closer approximation to in vivo condition. The hollow fiber bioreactor is a component of a hollow fiber cell culture system. A typical hollow fiber cell culture system, such as the CELLMAX™ 100 hollow fiber cell culture system (Cellco Advanced

Bioreactors, Inc., MD) contains a stendard glass medium bottle, which serves as the reservoir, stainless steel/Ryton gear pump, an autoclavable hollow fiber bioreactor, which contains the fibers and shell casing in which cells are cultured, and medical grade silicone rubber tubing, or other connecting means, which serves as a gas exchanger to maintain other connecting means, which serves as a gas exchanger to maintain the appropriate pH and pO₂ of the culture medium. All components are secured to a stainless steel tray of sufficiently small dimensions to enable four such systems to fit within a standard tissue culture incubator chamber. The pump speed and automatic reversal of flow direction are determined by an electronic control unit which is placed outside of the incubator and is generated to the gump motor via a flat ribbon cable which passes through the gasket of the incubator door. The pump motor is magnetically coupled to the pump and is lifted from the system prior to steam autoclaving.

The preferred HF bioreactor system for use herein is described in copending, allowed, U.S. application Serial No. 08/506,173; pages 1864-198

A HF system that closely emulates in vivo conditions thereby permitting T-cells to grow to densities of over 1 x 10³ cells/mls, preferably 1 x 10⁹ cells/ml, that uses fibers with a low molecular weight cutoff to retain mitogenic mAbs and serum components, and that does not have gradient formation problems, is described in copending, allowed, U.S. application Serial No. 08/506,173. This HF device allows outflow of the lumenal flow to be completely blocked. This leads to equal perfusion of nutrients along the entire length of the hollow fiber capillaries. It also includes an oxygen feed on the ECS of the bioreactor to provide desired oxygen delivery characteristics.

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ength of 14 inches, an ECS volume of volume of 120 mi, and a molecular weight cutoff (MWC) of 6,000 daltons were selected as the hollow fiber bioreactors for use in the hollow fiber processing apparatus.

To ensure equal distribution of nutrients across the entire length of these low.MWC cartridges, an automatic on/off solehold valve was placed on the outflow opening of the bioreactor. When the solehold is in the off position, medium is prevented from exiting the bioreactor. Instead, the medium ultrafiltrates to the cells in the ECS equally to all points of the bioreactor. The medium then passes out of the bioreactor through the ports. Ultrafiltration of nutrients is more physiological and therefore more desirable for maintenance of dense cultures of cells [see, e.g., Swaab st al. (1974) Cancer Res. 34:2814; and Davis et al. (1974) Cancer Res. 34:2814; and Davis et al. (1974) Chem. Eng.

All To remove the metabolic waste from the 'cells' in the 'ECS, the solenoid valve is switched to the 'con" position and the medium is "the medium to the controlled pressure to the ECS through the est protes. The medium then moves radially into the lumen. Finally, the medium is carried out the outflow opening.

The hollow fiber system permits the medium that ultrafiltrates from the lumen to the ECS (Cycle I) to be automatically replenished with oxygen and for the levels of glucose, lactate and carbon dioxide to be adjusted. This reconditioned medium is then returned to the ECS when the solenoid valve is opened in Cycle 2. The same adjustments are conducted for medium on the lumenal side of the bioreactor. In this manner, oxygen diffusion limitations can be overcome as oxygen is supplied to the lumen and the ECS of the bioreactor, eliminating diffusion across the hollow fiber capillaries as the sole means of oxygen transfer.

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For large-scale growth of regulatory immune cells hollow fiber bioreactors that have improved fluid dynamics to reduce gradient formation are preferable [see, e.g., U.S. Patent No. 4,804,628, see, especially, allowed copending U.S. application, Serial No. 08/506,173] are presently preferred. The hollow fiber bioreactors that have such improved fluid dynamics are best suited for the large-scale growth of regulatory immune cells.

In preferred embodiments, mitogenic monoclonal antibodies are coated onto the hollow fiber surafce in order to deliver the proper signals necessary to cause the immune cells to divide.

D. Effector cell expansion median production and the second secon

Effector cells are mononuclear cells that have the ability to directly eliminate pathogens or tumor cells. Such cells include, LAK cells, TiLs, CTLs and antibody-producing B cells and other such cells. These cells are produced by first treating cells collected from a patient in manner known to lead to differentiation of such cells. For example, TiL cells are produced by culturing solid tumor tissue obtained by blopsy in IL-2 and/or other agents that lead to TiL production. The cells are then activated and expanded in the presence of mitogenic agents, such as monoclonal antibodies specific for cell surface receptors or other agents, as described above for the regulatory cells.

In accord with the methods provided herein, the cells are not exposed to exogenous IL-2 (or any other agent upon which the cells will become dependent for in vivo activity or survival) and reinfusion is not accompanied by co-infusion of IL-2.

E. Selection of Immune Cell Phenotype and instrument of the second secon

Depending on the site of action at which a regulatory effect of infused cells is required for at which effector cells are required, different cell phenotypes may be required. Lymphocytes recirculate extensively.

throughout the body and then localize in tissues and lymphoid organs. This is accomplished by an array of adhesion molecules on lymphocytes and counter-receptors on the vascular endothelium, extracellular matrix and epithelium. Recent studies have identified several of the specific receptor/ligand interactions that mediate lymphocyte trafficking.

Infused cells that need to migrate out of circulation (e.g., to sites of inflammation) must have the capacity to move through extracellular matrix (ECM) of various compositions. For example, subendothelial basement membrane presents a barrier rich in type IV collagen, laminin and heparan sulfate proteoglycans. The ECM of the interstitium contains collagens I and III, as well as various glycosaminoglycans such as hyaluronic acid. Fibronectin and vitronectin are also encountered in basement membrane and interstitium. Immune cells can be loaded into columns containing these materials in order to screen for cells capable of migration through the interstitium.

It is also know that cells with a "memory" phenotype (i.e., CD45RA-, CD45RO+, CD29+, CD11a+, CD44+, CD54+, CD58+, L-selectin-) will accumulate non-specifically at sites of chronic inflammation. Cells that express L-selectin are least likely to migrate and should be used when the desired regulatory effect is required in the lymphatic organs.

Growing out cells with a defined antigen specificity may also be desired in order to prevent non-specific immunoregulation. Antigens should be selected that are unique to the site a regulatory effect is desired or to the disease-causing antigen(s).

F. Practice of the therapeutic methods

The therapeutic methods herein are designed to produce compositions containing clinically relevant [at least 10°, preferably 10¹0, cells or more] populations of regulatory immune cells and/or effector

immune cells for autologous infusion for treatment. The methods herein do not rely or use any agents for expansion that must be present after expansion to maintain cell viability or activity. In particular, expansion does not require or use IL-2. As a result, re-infusion of the cells does not require or use IL-2, thereby obviating toxicity and other problems associated with IL-2 infusion.

The compositions preferably contain substantially homogeneous populations of cells, such as Th1 cells or Th1-like cells, in which the cytokine profile is predominantly one type of cell (i.e., greater than about 10 50%). The compositions can contain regulatory immune cells, effector cells or both. In all instances the compositions contain clinically relevant, i.e., a therapeutically effective, numbers of cells.

Such compositions can be used therapeutically to restore an Immune cell imbalance. Immune cell imbalances are common in many 15 disease states. For example, a predominance of Th1 regulatory immune cells has been reported in autoimmune diseases such as rheumatoid arthritis [see, Simon, et al. (1994) Proc. Natl. Acad. Sci. U.S.A. 91:8562]; type I diabetes [see, Foulis, et al. (1991) J. Pathol. 165:97]; systemic inflammation [see, Brod, et al. (1991) J. Immunol. 147:810]: 20 inflammatory bowel syndrome [Niessner et al. (1995) Clin. Exp. Immunol. 101:428]; Grave's disease [see, de Carli, et al. (1993) J. Clin. Endocr. Metab. 77:1120]; Sjögren's syndrome [see, Oxholm, et al. (1992) Autoimmunity 12:185); primary systemic vasculitis [Grau (1990) Eur. Cytokine Netw. 1:203]; and rejected autografts [see, Benvenuto, et al. 25 (1991) Transplantation 51:887]. A predominance of Th2 regulatory immune cells has been reported in AIDS [see, Romagnani, et al. (1994) Res. Immunol. 145:611]; candidiasis [see, Puccetti, et al. (1995) Trends in Microbiology 3:237]; tuberculosis [Zhang, et al. (1995) Infect. Immun.

63:3231]; and allergy [see, Romagnani, et al. (1994) Curr. Opin. Immunol. 6:8381.

Also, the polarized Th1 and Th2 responses in humans to different antigens are known to play a role in protection, but also result in 5 immunopathology. The methods provided herein can be used to correct pathologic Th1 and Th2 responses by infusing autologous regulatory cells of the subset in short supply, thereby adjusting the ratios and absolute numbers. Since Th1 and Th2 cells have cross-regulatory properties: large infusions of the subset in short supply can counter-act the pathologic 10 effects of an imbalanced response. Some examples of the use of these methods and cells for treating several disease are provided. "It is the understood that the following are exemplary uses; any condition in which a pathologic T cell response is observed in which the ratios or amounts of particular subsets of T cells are outside the normal range can be treated by infusion of the T cell subset(s) that is in relatively short supply

The compositions of cell can be administered by any suitable means, including; but not limited to, intravenously, parenterally, or locally. The particular mode selected will depend upon the particular treatment and trafficking of the cells. Intravenous administration is presently preferred. Typically, about 1010-1011 cells can be administered in a volume of a 50 ml to 1 liter, preferably about 50 ml to 250 ml., more preferably about 50 ml to 150 ml, and most preferably about 100 ml. The volume will depend upon the disorder treated and the route of 25 adminstration. The cells may be administered in a single dose or in several doses over selected time intervals in order to titrate the dose. particularly when restoration of immune system balance is the goal.

1. Administration of agent and the Manager and

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Treatment of autoimmune disorders

macrophages, T-cells and B-cells.

The methods and composition of regulatory cell provided herein and a may be used to treat disorders that have an underlying autoimmune basis. or component.

a. Treatment of Rheumatoid Arthritis (RA)

RA is an immunologically mediated, chronic inflammatory disease characterized by synovial inflammation and autoantibodies. While the underlying cause of RA is unknown, it is well agreed upon that a fault in immune regulation is a principal factor contributing to the disease pathogenesis. Regulated control of normal immune responses are largely the result of interactions between, and the cytokine production of

AND TRAINING THE PROPERTY. Disease activity in RA patients has been positively correlated with the cytokine production of activated macrophages. In an inflamed joint, macrophages produce large amounts of pro-inflammatory cytokines which include IL-1, IL-6, IL-8, TNF-a and GM-CSF. These cytokines act to recruit Th1 memory cells to the joint and stimulate rheumatoid factor (RF) production leading to pannus formation and joint destruction. Treatment protocols which decrease the levels of proinflammatory Th1 cytokines in RA have been shown to result in clinical improvement.

The cytokines IL-4 and IL-10 are known to down-regulate macrophage activation and inhibit their production of IL-1, IL-6, IL-8 and TNF-q. IL-4 is also capable of suppressing the uncontrolled proliferation of synoviocytes, which is a major pathological feature of RA, IL-4 and IL-10 are produced by Th2 cells, which are virtually absent from the RA joint. Rather, RA joints have an abundance of Th1 cells."

Accordingly, RA can be treated by generating large numbers of the autologous, ex vivo derived Th2 cells from RA patients by the methods provided herein. The resulting cells, preferably in amounts greater than

10°, more preferably 10¹°, are re-infused into the patient to thereby suppress the chronic inflammatory lesions. Th2 cells of memory phenotype are preferred, since memory cells are most likely to migrate to the site of inflammation. In addition, the cells can be infused in an activated state; infiltrating T-cells in RA have been shown to have 5-6 fold increases in HLA-DR expression and 2-5 fold increases in VLA-T expression, both of which are activation markers.

It is also preferred that the infused Th2 cells only exert their regulatory action in the joints, so as to prevent a systemic immunosuppressive effect. Since the eliciting antigen is unknown in RA, the Th2 cells used should be specific for unique joint artifiens lear. Type II collagen or proteoglycani.

b. Treatment of Multiple Scienosis (MS)

MS is an autoimmune disease characterized by central nervous system inflammation and demyelination. The regulation of cytokine spectrum and production in MS is thought to have a decisive influence on disease outcome. Collective data has shown that Th1-associated cytokines, such as TNF-α, lymphotoxin, interleukin-12 and interferon-y promote disease, while cytokines from Th2 cells, such as IL-10, limit disease. In addition, TGF-β has been shown to be a disease downregulator. Studies in animal models of MS [experimental autoimmune encephalomyelitis (EAE)] have determined that a regulatory cell producing IL-10 and TGF-β, termed "Th3", has the greatest effect suppressing the development and inducing recovery from disease.

Accordingly, the methods herein can be used to generate therapeutic quantities of Th3 cells from MS patients for use in autologous cell therapy. Since recovery from disease is associated with infiltrating cells which produce IL-10 and TGF-\$\beta\$, the ex_vivo derived Th3 cells should preferably have a memory phenotype in order to enhance migration to the

inflammatory lesions. In addition, in order to make the immunosuppressive effect of the cells specific for the inflammatory lesions, cells specific for myelin or encephalitogenic epitopes of myelin antigens (e.g., myelin basic protein or proteolipid protein) should be used.

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IBD is a chronic inflammatory condition of the gastrointestinal tract. The etiology and pathogenesis of IBD is not known. Crohn's disease (CD) and ulcerative colitis (UC) are thought to be mediated by an abnormal or uncontrolled T-cell reaction to one or more common gut constituents. Active CD and UC are characterized by increases in Th1-like cytokines, with little to no detectable Th2-like cytokines.

Accordingly, the methods provided herein can be used to generate autologous. Th2 cells for infusion in IDB patients. Preferably, the infused cells will express the integrin, $a4.\beta7$: This integrin has been shown to be the ligand for mucosal addressin cell adhesion molecule-1 found on Peyer's patch high endothelial venules, which occur in the gastrointestinal tract. Lymphocytes which express $a4.\beta7$ will traffic to and are retained in mucosal organs. The gut mucosa is the site of chronic inflammation in IBD.

d. Treatment of Insulin-Dependent Diabetes Mellitus (IDDM)

IDDM results from the autoimmune destruction of pancreatic islet β cells by the host immune system. The destruction of islet cells is known to be mediated by T-cells. The NOD mouse is a spontaneous model of human IDDM. Islet transplantation as an isograft in these mice can produce normoglycemia and prevent and reverse early complications of diabetes. Host inflammatory responses, however, eventually lead to destruction of the islet transplants and disease recurrence. Analysis of

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these inflammatory responses has shown that graft specific Th1 cells mediate rejection, while Th2 cells are protective.

There is evidence that isograft and allograft rejection is madiated by Th1 cells and can be suppressed by Th2 cells. Th1 cells have been shown to actively promote-diabetes in NOD mice. Inhibition of Th1 cytokines leads to protection of silet isografts in NOD mice. Recently, it has been shown that the systemic administration of Th2 cytokines (IL²4 and IL-10) and adoptive transfer of an islet-specific Th3 cloine can inhibit syngeneic islet graft rejection in these animals. Furthermore, Th2-like responses have been shown to be protective in models of allogeneic organ and tissue transplantation.

Accordingly, the methods herein can be used to generate clinically relevant numbers of Th2 cells for infusion in IDDM patients that will protect against rejection of transplanted allogeneic siste cells. Preferably, the Th2-cells, will be specific for the allogeneic antigens on the transplanted islets. Alternatively Th2 cells specific for insulin can be used, Insulin-specific Th2 cells could also be used to treat fearly diagnosed IDDM patients to prevent islet destruction; as well as used in high risk patients as a vaccine to prevent or at least retard development of the diabetes.

e. Treatment of other autoimmune diseases

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Th1-mediated autoimmune diseases, such as, but not limited to, autoimmune thyroid diseases, anti-tubular basement membrane disease (kidney) Sjögren's syndrome, ankylosing spondylitis, ureoretinitis and others, can be treated by administration of compositions containing a clinically relevant, typically 10⁸-10¹¹, Th2 cells or a Th2-like composition.

the result in the part of the first parties that the

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3. Transplantation

permitting organ and tissue transplantation. For example, Th2 cytokines have been correlated with non-rejecting heart allografts, while Th1 cytokines correlate with rejection. The same is has been observed for renal allografts and mouse orthotopic liver allografts and skin allografts. Adoptively transferred Th2 cells suppress skin allograft rejection and also allow allogeneic engraftment of spleen cells in sublethally irradiated mice as well as suppress lethal GVHD (graft vs. host disease). T-cell mediated alloreactivity has been shown to be central in the pathogenesis of GVHD

Accordingly, the methods provided herein can be used to generate autologous Th2 cells for infusion in patients scheduled for organ or tissue transplant. Preferably, the Th2 cells will be specific for the alloantigens or an antigen unique to the organ or tissue being transplanted.

and graft rejection, was every laster that the second of the approximate to be a se-

4. Allergic Disorders.

Th2 cells appear to have a crucial role in initiating eosinophil infiltration which causes eczematous reactions in patients with atopic dermatitis, and airway hyper-responsiveness and pulmonary eosinophilia in allergic astima. Furthermore, atopic patients (patients with hayfever, dust and food allergies) have a preferential activation of Th2 cells. Recent evidence has shown that treatments that suppress Th2 development in vivo have profound inhibitory effects on allergen-induced airway changes and other atopic responses. Accordingly, since Th1 cytokines are known to inhibit Th2 responses, the methods herein can be used to generate large.numbers of autologous Th1 cells for infusion into atopic patients. Preferably, these cells will be specific for the allergen.

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5. Infectious Diseases and Cancer

An excess of Th2 cells is correlated with most infectious diseases, including viral, fungal, yeast, parasitic and mycobacterial infection. In order to change the regulatory balance in favor of cell-mediated immunity, Th1 cells could be infused into these patients. Prior art ACT protocols have used JlL and LAK effector cells and methods that use pathogen or tumor cell-specific CTLs. These effector cells would not be expected to work properly, in an immunocompromised host.

work properly in an immunocompromised host in the "help"

The co-infusion of Th1 regulatory cells should provide the "help"

10 necessary for the effector cells to perform their function and thus improve these therapies. Infusion of Th1 cells alone could provide sufficient help in vivo to drive endogenous CD8+ effector cells.

Accordingly, the methods herein could be used to generate large numbers of autologous Th1 cells for infusion into patients with infectious diseases or cancers. Preferably, the cells will be specific for antigens unique to the pathogen or tumor. The Th1 cells can also be infused with pathogen or tumor-specific cytolytic cells.

Of particular interest herein, are methods for treatment of HIV infection. Methods for producing virally purged CD4* cells are provided. In preferred embodiments, the cells are expanded under conditions in which Th1, cell differentiation is promoted. The resulting cells are reinfused into the donor HIV patient, whereby immunity will be restored. In other embodiments, these cells are reinfused with expanded effector cells, particularly effector cells that are specifically targeted against HIV infected cells.

compositions include, but are not limited to: influenza viruses, polio virus, leukemia viruses, hepatitis viruses, respiratory synctial virus, herpes viruses, retroviruses Epstein-Barr virus, syphillis (Treponema pallidum).

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cutaneous T-cell lymphoma (mycosis fungoides), Rhodococcus equi.
(intracellular respiratory pathogen), hypersensitivity pneumonitis,
onchocercal keratitis (river blindness), burn victims, chlamydia
trachomatis, mycobacterium avium, candida albicans, coxackievirus,
Leishmania major infection, cryptococcal infection and Bordetella

Infectious diseases that can be treated with Th2 cell compositions include, but are not limited to: filarial nematode (parasite), Plasmodium chaboudi (malaria), and Borrelia burgdofi (spriochete) infections.

Also of interest herein, are methods of treatment of cancer. In preferred embodiments, methods for treatment of renalcell carcinoma are provided. Transformed renal cells express heat shock protein hsp20. Consequently, hsp20-specific Th1 cells could serve as a cytokine delivery vehicle to increase local concentrations of IL-2 and IFNy in the tumor, thereby promoting anti-tumor effector cell function, activity and/or proliferation.

Th1 cells can also be used to mediate tumor regression in cancers including melanoma, breast cancer, head and neck cancer, prostate cancer and lung cancer. These is evidence that for certain tumors, a Th2 reponse may mediate regression.

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6. Vaccination

The development of effective vaccine strategies for intracellular pathogens, including, but not limited to, bacteria, viruses and parasites, is one of the major frontiers of medical research. Research centers on antigens from pathogenic organisms and adjuvants that can elicit a Th1-like response in patients. It is known that a Th1 response is protective for infectious pathogens. Th1 responses are weak or non-existent in some patients with most vaccine protocols. Other research focuses on eliciting an IqA antibody response, which is thought to be protective.

against organisms that enter the body through muscous membranes. An IgA response is mediated by Th2 cells. To better control the type of immune response a patient will elicit to a vaccine, the methods herein provide a means for <u>ex vivo</u> vaccination (<u>i.e.</u>, the addition of the vaccine antigen(s) to patient mononuclear cells <u>ex vivo</u>, whereby thecells are activated under conditions that promote the desired regulatory cell differentiation.

The methods provided herein can be used to withdraw blood from a patient, expose the isolated mononuclear cells to the vaccine antigen in 10 the presence of IL-12 and/or IFN-y and/or IL-4, and expand the Th1 or Th2 cells for reinfusion. Preferably, the cells used will have a memory phenotype so they will provide long-term protection. CD4+ and CD8+ Th1 or Th2 cells could be generated alone or in combination.

The following examples are included for illustrative purposes only

EXAMPLE 1

Screening mitogenic monoclonal antibodies

This example demonstrates a method for identifying antibodies that are suitable for expanding T-cell subsets, either singly or in combinations thereof.

In order to determine co-stimulatory signals required for T-cell subset proliferation, cells are incubated with various monoclonal antibodies (mAb) and their proliferation determined in ³H-thymidine incorporation assays. To exemplify this procedure, the following experiments were conducted.

Monoclonal Ab to CD3 (64.1, IgG2a) and anti-CD5 (10.2, IgG2a) were gifts from J. Ledbetter (Bristol Meyers, Seattle) and the mAb to CD28 (Kolt-2, IgG1) was a gift from K. Sagawa (Kurume University, Kyushu, Japan). These mAb were purified from ascites fluids on protein

A sepharose columns. All other mAbs were purchased from PharMingen (San Diego, CA). All mAbs were dialyzed against phosphate buffered saline and filtered through sterile 0.45 µm filters.

Goat anti-mouse affinity purified antibody. (Tago, Burlingame, CA)

was immobilized on plastic 96 well tissue culture plates. The antibody
was dissolved in sodium borate buffer (pH 8.6) at a concentration of
10 µg/ml and 100 µ was placed in each well. Plates were, washed three
times with RPME1-640 with 10% hormal human serum. Cells were
labelled with anti-CD3 mAb (1 µg/ml) on ice for 15 minutes prior to
10 plating. 50,000 cells were plated in each well. Co-stimulatory mAbs
were added in the soluble phase at 1 µg/ml. The cells were cultured at
37° C in an atmosphere of 5% CO₂. After 88 hours of culture, cells were
pulsed with 1 µCi of [PH]- thymidine (specific activity of 2 Cl/mole, New
England Nuclear). Eight hours leter, cells were harvested with a PHD cell
hervester (Cambridge Technology, Cambridge, MA) and the radioactivity
on the filter papers counted on a liquid scintillation counter (LS1701,
Beckman).

The results of mAb addition to purified CD4+ and CD8+ cells from a normal individual are shown below. Results are shown as mean counts per minute (cpm) of four replicates. Standard errors were always less than 10%.

Stimulation	CD4+	CD8+
medium alone	320	484
anti-CD3	582	541,
anti-CD3 + anti- CD5	18,450	17,222
anti-CD3 + anti- CD28	20,400	18,641
anti-CD5	450	246
anti-CD28	826	821

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These data demonstrate that anti-CD5 and CD28 are capable of providing a co-stimulatory signal for T-cell proliferation in CD4+ and CD8+ subsets when the cells are activated with anti-CD3. The results of combining anti-CD5 and CD28 are shown below.

5	Stimulation CD4+ CD8+	·
	medium	
	anti-CD3 585 500 508 11	
	anti-CD3+anti-CD5 13.422 10.080	
	anti-CD3 + anti-CD28	7
10	anti-CD3 + anti-CD5 + anti-CD28 25,248 29,804	_
	anti-CD3+IL-2 (10 U/ml) 11,428 12,401	Ì

These results show that the combination of anti-CD5 and anti-CD28 as co-stimulatory/signals in CD3 activated, purified T-cells induces
a greater proliferative response than either mAb alone. In addition, the
combined mAbs generated a proliferative response without addition of f

The effect of various mAbs (second signal) on purified CD8+ cells from a normal donor used in conjunction with anti-CD3 or anti-CD2 (first signal) was also tested. These results are shown below:

Stimulation	aCD3	aCD2	Medium
aCD5	. 206 _	193	155
aCD8	787	578	640
αCD11a_	949	830	.840
αCD27	844	Ż	788
aCD28	1928	529	640
aCD44	779	477	498
aCD45RO	3199	1878	1978
IL-2	4347	1834	nd
Medium	289	217	212

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These results demonstrate that anti-CD3 as the first signal delivers a more powerful proliferative stimulus than anti-CD2. Anti-CD45RO and anti-CD28 mAbs appear to deliver the strongest second or co-stimulatory signals when used with anti-CD3.

Combinations of these antibodies were tested on anti-CD3:
activated, ex vivo generated CD8+ cytolytic cells specific for the
MAGE: 3 antigen on melanoma cells. These results are shown below:

ing the st	anti- CD11a	anti-CD2	7 anti-CD2	28 anti- Sw vi. v CD45RO
anti-CD11a		1365	1116	1208
anti-CD27	1365	14.74	374	973
anti-CD28****	1116	374	100.00	948
anti-CD45RO	665	973	948	

Combinations including anti-CD11a provided the strongest proliferative signals for these cells. None of these combinations provided very exceptional growth. This sometimes occurs in CD8+ CTL/which are unable to produce sufficient endogenous cytokines. Co-culturing of these cells with autologous CD4+, however, enhanced the proliferation of these cells with mAb stimulation. This probably resulted from the increased endogenous production of ill-2, as well as IFN-y and II-7.

EXAMPLE 2

CD4+and CD8+ T-cells from Normal Donor

This example demonstrates that polyclonally activated CD4+ and CD8+ regulatory T-cell subsets can be expanded without IL-2 to clinically relevant numbers from a starting number of about 1 x 10⁶ cells using the disclosed methods.

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A. Collecting mononuclear cells

Mononuclear cells from normal donors were obtained from source leukocyte packs (Interstate Blood Bank, Inc.). The leukopack cells were diluted 1:1 with Hank's Buffered Salt Solution (HBSS) without calcium (Ca²⁺) or magnesium (Mg²⁺) and 30 to 35 ml of the diluted cells were placed over 12 ml of Ficoll-Hypaque and the tube centrifuged at 1500 RPM at room temperature: The buffy coat layer containing lymphocytes and monocytes was transferred by Pasteur pipette to a clean 50 ml centrifuge tube and washed three times with HBSS. The cells were then resuspended in RPMI-1640 medium supplemented with 10% human serum, 25 mM HEPES buffer, 2.0 mM glutamine; 1.0 mM sodium pyruvate, 0.1 mM non-essential amino Acids, 2 x 10° M,2-mercaptoethanol, 10 IU of penicillin G and 100 mg/ml streptomycin sulfate (cRPMI). The monocytes were depleted by adherence to plastic T-cell flasks incubated overnight at 37°C in an amosphere of 5% CO₂ and 100% humidity.

T-cell subsets were purified with immunomagnetic bead technology. GAM-coated beads (Dynal, Inc.) were washed twice with HBSS and incubated overnight on a rotating wheel at 4°C in HBSS with 1% normal human serum in order to block nonspecific binding. The non-adherent cells were incubated with either anti-CD4 or anti-CD8 mAb at pre-titered concentrations on ice for 30 minutes. Labelled cells were washed twice and resuspended in cRPMI at 10 cells/ml: The beads were added to the cells at a bead/cell ratio of 2:1 and mixed well. This mixture was gently centrifuged at 500 RPM for 1 minute at 4°C. The bead/cell mixture was then placed on a rotating wheel for 30 minutes at 4°C. The bead/cell mixture was then diluted 5 fold with cRPMI and placed on a

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cobalt salarium magnet. The supernatant was aspirated and rosetted and the procedure repeated. The rosettes were incubated for 24 hours in cRPMI at 37°C in an atmosphere of 5% CO₂. After 24 hours, the majority of cells detached from the beads and the beads were removed by placing the solution back on the magnet. The resulting cells were greater that 98% pure CD4⁺ or CD8⁺ T-cells as assessed by flow cytometry.

C. Ex Vivo Differentiation

The purified CD4+ cells were divided into twosparate groups of 1 million cells each. The first group was activated with immobilized anti-CD3 mAb in the presence of 400 U/ml of IL-4 and 10 µg/ml of anti-IFN+ mAb and anti-CD28 mAb. This first group (Th2) was expanded under these conditions for another 10 days. The second group was activated with immobilized anti-CD3 in the presence of 25 U/ml of IL-12 and 150 U/ml of IFN-y, and anti-CD28 mAb. These cells were harvested and washed after 6 days of culture.

D. Regulatory cell expansion.

One million of each of the purified T-cell subsets were labelled for 30 minutes on ice with anti-CD3 mAb (64.1, IgG2a). 2.5.X 10⁵ cells of the purified CD4⁺ and CD8⁺ cells were suspended in 1 ml of-cRPMI and plated into 4 separate wells of a 24-well plate coated with goat anti-mouse (GAM) polyclonal antibody. Purified anti-CD5 (10.2, IgG2a) and anti-CD28 (KOLT-2, IgG1) mAb were added to the wells at a final concentrations of 200 ng/ml. The cells were then incubated at 37°C in an atmosphere of 5% CO₂.

After 3 days, 1. ml of cRPMI with 200 ng/ml of anti-CD5 and anti-CD28 was added to the wells. After 6 days, the wells were harvested, pooled and washed twice in cRPMI. The viable cells were counted and resuspended in cRPMI at 1 x 10⁶ cells/ml and incubated in T-flasks for 48

hours at 37°C. The cells were then harvested, washed twice, labelled with anti-CD3 mAb on ice for 30 minutes and inoculated into the extra capillary space of a GAM-coated mini-hollow fiber bioreactor with 200 mg/ml of anti-CD28 an danti-CD5 mAb. The cells were harvested, washed and counted after 14 days.

1. Mini-Hollow Fiber Bioreactor

A mini-hollow fiber device was constructed to expand immune effector cells. The device had four mini-hollow fiber units in parallel. The hollow fibers (CD Medical, Hialeah; FL) had a 9 ml extracapillary volume 10 and the fibers had molecular weight cut offs of 10,000 dations. The hollow fibers were coated with GAM polyclonal antibody. Coating was accomplished by dissolving GAM polyclonal antibody, at a concentration of 10 mg/ml, in sodium borate buffer (pH 8.6) and inoculating the sterile solution into the extracapillary space (ECS) of the hollow fiber 15 bioreactors. The lumenal and ECS ports were then sealed and the bioreactors placed on a rotating plate and incubated at 4°C for 24 hours. Prior to use, the bioreactors were washed with phosphate buffered saline with 1% normal human serum.

The flow path included an integration vessel, pump and oxygenation cartridge. Luminal flow rates ranged between 100 and 400 ml/minute and were increased manually proportionate with the cell growth in the bioreactors. The pH and temperature were continually monitored and controlled by microprocessor. The pH was adjusted and maintained at 7.2 by altering the speed of fresh medium fed into the integration vessel and the percent CO₂ in the oxygenation cartridge. The temperature was controlled to 37°C by adjusting the wattage to a heating coil wrapped around the integration vessel.

2. Single Large Hollow Fiber Bioreactor

The cells recovered from the mini hollow fiber device were incubated in T-flasks at 1 x 107 cells/ml in cRPMI without mAb stimulation for 48 hours. The cells were then labelled with anti-CD3 mAb and inoculated into a GAM-coated large hollow fiber bioreactor [see. copending allowed U.S. application Serial No. 08/506:173 discussed above] with 200 ng/ml of anti-CD5 and anti-CD28 mAb. The cells were harvested, washed and counted after 14 days, now a 400 harvested

8-Cartridge Hollow Fiber Bioreactor

The cells recovered from the single large hollow fiber bioreactor [see, copending allowed U.S. application Serial No. 08/506:173. discussed above) were incubated for 48 hours in a 10 liter spinner flask at 107 cells/ml in cRPMI without mAb stimulation. The cells were then labelled with anti-CD3 mAb and inoculated into each of the 8 GAM-15 coated hollow fiber bioreactors with 200 ng/ml of anti-CD5 and anti-CD28 mAb... After 14 days, the cells were harvested, washed and counted. The second of the country of the contest of the country of the contest of the country o

E. Results of the second second second and the first terms.

Clinically relevant numbers of cells were produced as follows:

1	Day	CD4* (Th1)	CD4 * (Th2)	CD8+	Culture Vessel
1	CO	1x10 ⁶ cells	1x10 ⁶ cells	1x10 ⁶ cells	24-well plate
١	- 6	1.3x10 ⁷ cells	7.2x10 ⁶ cells	9.8x10 ⁶ cells	24-well plate
1	80 1	1.0x10 ⁷ cells	6.5x10 ⁶ cells	6x 10 ⁶ cells	Mini-HF
1	22	1.3x10 ⁹ cells	1.0x10 ⁹ cells	1.2x10° cells	Mini-HF
I	.: 24	1.1x10° cells	1.0x10° cells	1.1x10° cells	1-large HF
l	.38	1.4x10 ¹⁰ cells	1.0x1010 cells	1.2x1010 cells	1-large HF
l	40	1.3x10 ¹⁰ cells	1.0x1010 cells	1.0x1010 cells	8-Large HF
ı	54	1.1x1011 cells	1.0x1011 cells	9.9x1010 cells	8-Large HF

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anti-CD28 mAb.

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Therefore, compositions containing clinically relevant numbers of Tretaining a sequence of the second cell subsets can be produced.

EXAMPLE 3

Virus-purged CD4* Th1-cells from HIV* patient

- This example demonstrates that clinically relevant numbers of virus-purged CD4+ Th1-cells can be generated by the methods herein for use as an ACT for A.I.D.S. The cells were purged of active virus by selection of CD4 antigen and were polyclonally activated and again selected for CD4 antigen to purge of latent virus.
- A. Obtaining Mononuclear Cells An HIV+ patient, identified by a routine blood screening procedure confirmed by Western Blot analysis, in WHO stage IV was the donor for this study. The patient underwent a leukopheresis procedure for collection of peripheral blood mononuclear cells.
- B. . . Regulatory cell purification CD4+ cells were isolated by positive selection on immunomagnetic beads as described above. The CD4+ cells were then activated in the country of th 24-well plates with immobilized anti-CD3 mAb and in the presence of 40 U/ml of interferon-y (IFN-y). After 24 hours in culture, the cells were 20 harvested, washed and re-selected for CD4 on immunomagnetic beads. The positively-selected cells were labelled with anti-CD3 mAb and plated at 25.000 cells/well in a GAM-coated 96-well plate in cRPMI. Anti-CD28 mAb and IFN-y was added to the wells at a concentration of 1 µg/ml and 40 U/ml, respectively. After 7 days, supernatant from each well was tested for p24 antigen with a commercial ELISA assay (Dupont). All negative wells were pooled, relabelled with anti-CD3 mAb and replated at 25,000 cells/well in a GAM-coated 96-well plate in cRPMI with

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C. Regulatory cell expansion

The cells were expanded as described in Example 2 above, except that only anti-CD28 mAb was used as a co-stimulatory agent.

6.3 x.1010 cells were grown over a 72 day period. The cells were negative for p24 antigen and were capable of producing IL-2 and IFN-y, but little or no IL-4. The cells were also shown to be capable of providing help for NK function in a dose-dependent manner. The cells were reinfused into the patient. Reinfusion of these cells into the HIV* patient should be a treatment for A.I.D.S.

EXAMPLE 4

HIV-specific CD8* cells from a HIV* donor

This example demonstrates that antigen-specific CTL can be purified and expanded from an individual with a viral infection.

A. Obtaining Effector Cells

3 x 10° mononuclear cells were obtained by leukaphoresis from a stage IV A.I.D.S. patient., CD8*; CD25* cells were purified by two rounds of selection on immunomagnetic beads.

B. Expansion of Effector Cells

Approximately 2 x 1.0⁸ cells were recovered and expanded in a 24-well plate coated with anti-CD3 mAb and with soluble anti-CD28 mAb. After 6 days, the cells were washed (x 2) and inoculated into mini-hollow fiber bioreactors. After 18 days in the mini-hollow fiber units, the cells were washed, counted and allowed to rest 2 days before inoculation into a cartridge of the large hollow fiber bioreactor under the same conditions as described in Example 2 above.

After 16 days, the cells were harvested, washed and allowed to rest for 2 days. The viable cells were then inoculated into the 8-cartridge

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hollow fiber bioreactor system and cultured under the same conditions as described in example 2 above.

C. Results

6 x 10¹⁰ viable cells were harvested after 20 days. The cells showed significant Ag-specific CTL activity against infected autologous cells.

These cells can be reinfused into the patient as a treatment for A.I.D.S. In addition, these can be co-infused with virally-purged CD4+, produced as described in EXAMPLE 3.

EXAMPLE 5

relevant numbers.

Antigen-specific Th2-like cells from a normal donor

This example demonstrates that antigen-specific Th2-like CD4+
cells can be derived from a normal individual and expended to clinically

A. Obtaining regulatory Cells

50 ml of blood was collected into a heparinized syringe, using sterile technique, from an HIV volunteer. Peripheral blood mononuclear cells (PBMC) were separated by Ficoll-Hypaque density gradient centrifugation. The PBMC were cultured in 10 ml T-flasks at 2 x 10° cells/ml and pulsed with gp120 antigen in cRPMI that contained 1.0 µg/ml of anti-FN-y mAb and 20 U/ml of IL-4. After 2 days, the blasts were collected by selection of CD25 on immunomagnetic beads. The blasts were allowed to rest for 72 hours and were than re-stimulated with gp-120 pulsed, autologous monocytes and immediately cloned in soft agar. The small number of cells that survived and grew out as colonies (1/150,000) were enriched in Ag-specific cells that produced IL-4 and IL-10 and little IFN-y upon stimulation, and, thus, were Th2-like in cytokine profile.

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B. Expansion of Effector Cells

The cells were expanded as described in Example 2 and grew to 9×10^{10} cells in 62 days.

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Differentiation of Th2 cells from Precursors in Rheumatold Arthritis
Peripheral Blood

While T cell cytokine expression is very low in rheumatoid arthritis
(RA), the absence of Th2 factors (e.g., IL-4 and IL-13):is especially
striking. Since Th2 cytokines suppress production of pro-inflammatory
cytokines, metalloproteinases and rheumatoid factor, their relative
absence in RA could contribute to disease perpetuation. The lack of Th2
cells in synovium suggests that this differentiation pathway might be
defective in RA. To determine if Th2 precursors are present in RA, the
ability of peripheral blood RA CD4 + T cells to differentiate into Th0 (IL-4
15 + IFN-4), Th1 (IFN-4, no IL-4) and Th2 cells (IL-4, no IFN-4) in vitro was
studied.

Purified CD4.+ T cells were cultured in the presence of immobilized a CD3 antibody, alL-12 and IL-4 for 3 d. Cells were then washed and stimulated with PMA and ionomycin in the presence of monerain for 6 hr.

The cytokine phenotype was determined using 2-color flow cytometry on permeabilized cells with alL-4 and BIFN-4 monoclonal antibodies. The results are shown as percent cells ± standard error (se); "n" values are in parentheses.

1 34	Treatment	Th2(%)	Th0(%)	.Th1(%)
RA (9)	øCD3	0.68±0.19	0.44±0.11····	10.38 ± 2.61
Normal (6)		0.56±0.08	0.55±0.17	11.07 ± 2.89
RA (4)	aCD2+IL-4	1.43±0.32	0.29±0.09	4.68±0.91
Normal (5)		1.50±0.26	1.69±0.56	13.27±2.46
RA (6)	aCD3 + alL-12 + IL-4	3.03±0.92'	1.68 ±0.44	12.51 ± 3.15
Normal (3)		1.45±0.35'	0.72 ±0.36	7.30±0.84

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These data indicate that similar numbers of Th2 cell precursors are present in the peripheral blood of normals and patients with RA.

Furthermore, the mature Th2 cell population can be significantly increased (p<0.05) with IL-4 and o-IL-12 antibody. Hence, a specific Th2 precursor defect does not account for the cytokine profile in the joint. This raises the possibility that novel therapeutics could be developed involving the administration of ex vivo differentiated and expanded Th2 cells.

eura de aprez de la del EXAMPLE 7 de

HIV +: Lymphocyte Proliferation: a few and the second few and the seco

The ability of PBL from HIV + donors to proliferate in response to the polyclonal activator PHA-P and immobilized anti-CD3 mAb was compared with PBL from a normal donor (Table 11). PBL from HIV + donors exhibited a marked suppression in the ability to respond to either mitogenic signals when compared to PBL from normal donors.

Table 1: Comparison of Proliferative Response of Normal

PBL Source	Medium Alone	PHA-P (1 ng/ml)	Immobilized anti-CD3 mAb
normal donors	1,446 ± 241	25,813±1200	27,206 ± 1891
HIV + donors	2,041 ± 421	5,680 ± 460	4,204 ± 562

Peripheral blood lymphocytes (PBL) isolated over Ficoil-Hypaque were plated at 50,000 cells/well in 96-well flat bottom culture plates. Cells were pulsed after 88 hours of stimulation with medium alone, PHAP or immobilized anti-CD3 mAb with [3H]-thymidine for eight hours and the average mean and standard error of quadruplicate samples for six normal and six HIV + individuals is shown in cpm.

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To determine if purified T-cell subsets from HIV+ donors were capable of responding to mitogenic stimuli in the absence of activator, the following study was conducted. PBL from six normal and six HIV+ individuals, isame individuals, as used in the experiments shown in Table 1) were incubated in plastic fissue culture dishes for 24 hours at 37° C in an atmosphere of five percent CO₂ In air. The CD4+ and CD8+ T-cell subsets were purified using positive selection on immunormagnetic

beads as described previously. The results are shown in: Table 2. Proliferative Response of Normal and HIV+ T-Cell Subsets to accommod to the control of the

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Immobilized anti-(purity %)CD4* Medium **PMA** CD3*IL-2 (99.5) Normal donors 1.841 ± 320 42,186 ± 3444 35,920±3420 1.346 ± 230 29.212±1841 31,440 ± 6210 (98.8) HIV + donors (purity %) CD8++ 1:925 ± 421 12.420 ± 821 10.920 ± 1104 (98.8) Normal donors (98.4) HIV + donors 1.212 ± 168 10.861 ± 948 6.155 ±.718

T-cell subsets isolated by positive selection on immunomagnetic beads from six normal and st NIV.+ donors. Average purities are shown in parenthesis, The cells were plated at 50,000 cells per widel in 36 well flat bottom itsus culture plates in CRPMI and 10 percent NHS pulsed for eight hours with 1 µCl [PH] - thymidine after 88 hours of stimulation with either medium alone, immobilized anti-CD3+ II.-2 (10 µml) or PMA (0.5 ng/ml). Results are shown as the average cpm_and standard errors, Each group was performed in triplicates.

The results indicate that a significant T-cell proliferative response is possible from HIV+ donors. The CD4+ cell response to anti-CD3+ IL-2 of HIV+ donor cells was approximately 30 percent less than for the normal donors, but still significantly higher than the medium alone control. The CD8+ cells of HIV+ donors responded nearly the same to anti-CD3+/IL-2 as did normal cells. The CD8+ response of normal and HIV+ donor cells was significantly less than that observed in CD4+

transmitte was set

cells. These results indicate that purified T-cell subsets from HIV+

To demonstrate that mitogenic mAbs could provide the second signal for T-cell proliferation in anti-CD3 activated T-cells from HIV+ donors the following experiments were performed. T-cells purified from PBL of HIV+ donors were isolated using AET-treated SRBC. The anti-CD3 activated T-cells were exposed to soluble anti-CD8 alone, anti-CD5 alone and a combination of anti-CD28 and anti-CD5. The results are shown in Table 33

Table 3 Proliferation Response of T-Cells from HIV+ Donors

Stimulation	cpm ± SEM
medium	1,810±130
anti-CD3	2,338 ± 144
anti-CD3±rIL-2	/se ::11;882±35 ===
anti-CD3± anti-CD28	13,334±300
anti-CD3 ± anti-CD5	3,629 ± 102
anti-CD3 ± anti-CD5 + anti-CD28	12,882±69

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T-cells purified by the AET-treated SRBC E-rosetting procedure (99.6 percent CD3+1 were isolated from PBL of an HIV+ donor. The cells were plated at 50,000 per well in a 96 well flat bottom tissue culture plate in cRPMI and 1.0 percent. NHS. The cells were activated with immobilized anti-CD3 mAb and stimulated with either IL-2 (10 u/ml), soluble anti-CD28 mAb (200 ng/ml) soluble anti-CD5 (200 ng/ml) or a combination of soluble, anti-CD8, and anti-CD5. Cells were pulsed for eight hours with 1 u Ci [*H]- thymidine after 88 hours of stimulation. Results are shown as cpm and standard error from a single donor. Each treatment group was run in guadruplicate.

Anti-CD28 was as effective as IL-2 in providing the second signal to purified T-cells from an HIV+ donor, Anti-CD5 had no effect alone or

in combination with anti-CD28 while augmenting the proliferative response in T-cells from normal donors.

Minimum Cell Density Required for Proliferative Response.

In order to determine the minimum cell density required for the simmobilized anti-CD3/soluble anti-CD28 system to cause 7-cells from HIV + donors to proliferate, the following study was conducted.

T-cells from an HIV+ donor and a normal donor were purified using the AET-treated SRBC E-rosette procedure described earlier. Purities of T-cells were 99.4 pergent for the HIV+, donor and 99.2 percent for the normal donor. The T-cells were serially diluted from a starting concentration of 1 × 10° cells/ml and plated onto 96 well plates. Final cell country/well ranged from 100,000 to 1,000. All experimental groups were

Table 4. Minimum Cell Density Required for T-Cell Proliferative Response in the Anti-CD3/Anti-CD28 System

studied in quadruplicate. The results are shown in Table 4.

rantidatili on at VIP to acree. earn was Normal Donor HIV + Donor # Cells/Well Medium . . Anti-CD3 : Medium Anti-CD3 Anti-CD28 Anti-CD28 20 100,000 1.628±42 22.842 + 462 1,042 ± 214 52.820±428 50,000 1.822 ± 120 14.920 + 1081,944 ± 108 29.642 ± 262 25,000 1.206 ± 24 8,444 ± 48 1,496±51 14.322 ± 125 10,000 1,828±18 2,420 ± 186 1.684±49 6.246 ± 68 5,000 1.484 ± 56 1.848 ± 342 1.544 + 324.820+320 25 1.296 ± 260 1.000 1,741 ± 85 1.821 ± 74 1.948 + 146

T-cells purified by an E-rosetting procedure using AET-treated SRBC from a normal and an HIV+donor were tested for their ability to respond to immobilized anti-CD3 mAb and 200 ng/ml of soluble anti-CD28 mAb. T-cells, were cultured for 88 hours with anti-CD3/anti-CD28 or medium alone and then pulsed with IPH- triymidine for an additional eight hours. Results are shown as cpm ± standard error. All treatment groups were run in duplicate. A single donor was used in each treatment group.

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T-cells from the HIV+ donor exhibited significant proliferative response in the anti-CD3/anti-CD28 system at cell densities above 2.5 x 10⁸ cells/ml (25,000 cells per well). T-cells from the normal donor were capable of responding down to a density of 5 x 10⁸ cells/ml (5,000 cells/well). The proliferative response of T-cells from the HIV+ donor was approximately 50 percent less than the T-cells from the normal donor.

HIV Purge Method 10 to the last term of the black to the term of the contract to the contract

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H9 Continuous Cell Line. In order to reconstitute the Immune system of an AIDS patient; large numbers of CD4 + cells are required. Since these cells harbor latent and active HIV-1, a method is required that will isolate a viral-free starting population of CD4 + cells. If the purging method is not 100 percent effective; the virus will quickly take over the culture as it is stimulated to replicate by activation of the host cell.

To demonstrate the feasibility of purging CD4+ cells from AIDS patients of HIV-1, an HIV-infected continuous cell line was used. The cell line, H9 (gift from Dr. Gallo, NIH, deposited under ATCC No. CRL 8543), is a cloned CD4+ human lymphocyte line. H grows continuously in culture and can also continuously propagate HIV-1.

p24 ELISA. A commercial kit (Dupont) was used to assay the amount of virus in the cell cultures and monitor the efficiency of the purging experiments. The kit can detect one viral particle in 5,000 cells. The test uses highly specific rabbit polyclonal antibodies to HIV p24 core antigen. These antibodies are immobilized on a 96-well plate. The antibodies capture p24 antigen that is released into the supernatant of a cell culture after treatment with five percent triton-X to lyse the cells. The captured p24 core antigen is then complexed with anti-p24 biotinylated polyclonal antibodies. The complexes are probed with a streptavidin-HRP (horseradish peroxidase) conjugate. The complexes are

detected by incubation with orthophenyldiamine-HCI (ORD) which produces a yellowish color proportional to the amount of HIV p24 antigen captured. The absorbance of each well was determined on a microplate reader (Dynatech, Minireader II) and calibrated against the absorbance of known values of p24 antigen. To increase the sensitivity of the test, test cells were co-cultured with PHA-activated, normal lymphocytes.

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phenotypic behavior of infected cells. HIV+ cells with active virus will express the env gene products gp120 and gp41 on their cell surfaces.

Since it was reported that HIV+ cells with active virus internalize their CD4 receptors, positive selection of CD4 was tested.

whereas infected-H9 cells (H9 +) are four percent CD4+ (H9-) second to by flow cytometry. An experiment was designed where 10 million H9 cells were mixed in the following ratios:

(1) and 00 percent H9 + and 70 percent H9-;

(3) 60 percent H9 + and 30 percent H9-; and

(3) 60 percent H9+ and 30 percent H9; and
20 (4) 80 percent H9+ and 20 percent H9

Cells from each group were positively selected for CD4 with

Impunomagnetic beads. A sample of the positively selected cells were

immunomagnetic beads. A sample of the positively selected cells were tested for p24 with the commercial ELISA test (no co-cultivation).

Results are shown in Table 5.

Table 5 Purge of H9 Cells Infected with HIV-1

gir av a med j	p24 before CD4 removal	
0%H9+	0.03 ng	0.01 ng
10%H9+	0.25 ng	0.00 ng
30%H9+	0.58 ng	0.00 ng

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	p24 before CD4 removal	p24 after CD4 removal
60%H9+	0.94 ng	0.03 ng*
80%H9+	1.36 ng	0.03 ng*
100%H9+	2.14 ng	0.09 ng

5 *same as negative control

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The continuous cell line H9 infected HIV-1 (H9+) and non-infected H9 (H9-) were mixed at various ratios. Cells expressing the CD4 surface antigen were purged from the mixture using specific mAbs and immunomagnetic beads. The amount of p24 antigen in the cultures was determined before and after the purge process.

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All groups with the exception of the 100 percent H9+ group were successfully purged of virus below the detectable limits of this assay. To determine if the negative fractions would continue to be viral-free the cells were incubated for 20 days in 24 well plates with 3 x 106 indicator cells (normal lymphocytes activated with PHA for 72 hours) in cRPMI and 10⁸ NHS. Fresh indicator cell were added again on day seven. On days seven, 14 and 20, 1 x 108 cells from each group were lysed with triton-X and assayed for p24. The results are shown in Table 6.

20 Table 6. Co-Cultivation of Viral Purged H9 Cells with Indicator Cells

-	THE RESERVE OF THE PERSON NAMED IN			
Day	10% H9+	30% H9+	60% H9+	80% H9+
0	0.00 ng	0.00 ng ;	0.03 ng	0.03 ng : : :-",
7	0.04 ng	0.14 ng	0.20 ng	0.29 ng
14	0.09 ng	0.23 ng	0.38 ng	0.32 ng
20	0.25 ng	0.53 ng	0.59 ng :	0.38 ng

H9+ cells mixed with H9-cells at various ratios were pluged of CD4 - cells using immunomagnetic baseds. The H9-fractions were co-cultured with PHA-stimulated lymphocytes. The fractions were tested for presence of p24-viral antigen at days zero, seven, 14 and 20.

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These results indicate that the original viral purge was not 100 percent effective and virus can still exist below the level of sensitivity of the assay. In a further attempt to develop a viral-free culture, 1 x 10° cells from each group were serially diluted and plated at 500 cells per well in 2,000 wells of 24-well plates. The cells were allowed to expand for 14 days and then were co-cultured with indicator cells for 20 days as before. Cell samples were analyzed for p24 antigen after 20 days as described earlier. The results are shown in Table 7.

Table 7. Co-Culture of Viral-Purged H9 Cells with Indicator Cells
After Plating at 500 Cells/Well

Group	% of Positive Wells*
10%н9+	16%
30%H9+	32%
60%H9+	26%
80%H9+	32.5%

^{*}any value over the negative control

40 H9+ cells mixed with H9- cells at various ratios, purged of CD4+ cells and cultured for 20 days with PHA-stimulated indicator lymphocytes were serially diluted to 500 cells per well of a 24-well plate. The cells were allowed to expand for 14-days and assayed for p24 viral antigen. The percent of wells from each ratio of H9+ to H9- cells that were positive for p24 is shown.

Those results showed that virally-infected cells could be eliminated after positive selection by serial dilution. To further validate this procedure, the negative wells were pooled and cultured with indicator cells for another 20 days. All groups remained negative for p24 antigen (data not shown). Thus, the combination of positively selecting CD4+cells followed by serial dilution, should be useful as a viral purge method.

To further test the sensitivity of the assay system, two-fold serial and dilutions were made from H9+ cells from 500 cells/well to less than one

cell/well (defined as a two-fold dilution beyond one cell/well). The results are shown in Table 8.

Table 8. Serial Dilution of H9+ Cells to Test Sensitivity of p24 Antigen

5	Positive Control		ு ந். H9 + ,Cells ு ் பிறிந்	
	Concentration ng/ml	Absorbance	Concentration	Absorbance
	0.25	1.03	> 8 cells/well	- ";over ;:
	0.125	0.55	8 cells/well	1.53
	0.0625	0.30	4 cells/well	0.89
10	0.0313	0.15	2 cells/well	0.53
	0.0157	0.04	1 cell/well	0.24
	0.0 ng/ml	0.03	< 1 cell/well	0.10

Absorbance of known concentrations of p24 antigen in a commercial ELISA

(Dupont) were compared with absorbance of cell lysates from an HIV-1 infected continuous cell line - H9.

These results indicate that the assay is extremely sensitive; it is able to detect p24 in < one cell/well down to 0.0157 ng/ml concentration.

20 Viral Purge from HIV + Donor

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The H9 studies indicated that positive selection of CD4+ cells combined with serial dilution could isolate a viral-free subpopulation of cells. The process can be monitored with great sensitivity by a commercial p24 assay. This process, however, does not address the purging of latent virus from the cells. In order for latent virus to proliferate, the host cell must be activated. The immobilized anti-CD3 system has proven to be an effective activator of these cells. After activation, the viral-free cells must be protected or they will soon become infected just as the indicator cells do in the p24 assay. Anti-CD4 mAb

30 was used to protect uninfected CD4+ cells.

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Materials and Methods

Lymphocytes were Isolated from the AIDS patient following leukaphoresis as described above. A sample of unfractionated cells were tested for p24 in a co-cultivation test for 20 days. Similar samples were tested after macrophage adherence, CD4 positive/selection and CD8 positive/selected for with GAM-coated immunomagnetic beads. The positively selected cells were relabelled with anti-CD3 and placed on GAM-coated 96-well plates at 25,000 cells/well. Anti-CD28 was added to the growth medium.

After seven days, supernatant from each well was tested for p24 antigen. All the negative wells were pooled and again subjected to CD4 positive selection with immunomagnetic beads. The positively selected cells were relabelled with anti-CD3 mAb and plated again at 25,000 cells per well. Anti-CD28 was added to the medium and the wells were tested for p24 again after seven days. Negative wells were again pooled and expanded as described previously for normal lymphocytes with the exception of only anti-CD28 and the addition of anti-CD4 (leu 3a; Becton Dickinson) to protect the cells from any residual virus. The cells were expanded to over ten million and a one-million cell aliquot was harvested for co-cultivation with indicator cells, p24 readings of cell lysate was taken after 20 days.

25 Results

Results are shown in Table 9.

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Table 9 Viral-Purge of Lymphocytes from HIV + Donor.

p24 Levels	\$7.410 0
PBL (before adherence)	0.32 ng
PBL (after adherence)	0.28 ng
CD4 +10/304 District of 1990 4 in	0.24 ng
CD8+ at long trace!	0.00 ng

Amount of p24 antigen recovered from a one million cell yeate of HIV+ cells before removal of macrophages by adherence to plastic T-flasks; after the removal of macrophages, after positive selection of CD4+ cells and CD8+ cells.

The CD4+ cells were plated at 25,000 cells per well of a 96-well plate and expanded for seven days on immobilized anti-CD3 mAb and soluble anti-CD28 mAb. Each well was then assayed for p24 antigen.

Results are shown in Table 10.

Table 10 Detection of HIV-1 in Wells of Expanded

11 July 1	# of Wells	# Greater than Background	% Negative
Group 1	133	. 24	82%
Group 2	108	18	83%
Group 3	141	. 29	79%

Amount of p24 antigen recovered from wells of 96 well plates with 25,000 CD4+ cells purified from the peripheral blood of an AIDS patient and expanded for seven days on immobilized anti-CD3 mAb- and sotuble anti-CD28 mAb. Each group represents the results of a separate purification from the same patient.

The percent negative wells was very consistent. The cells from the negative wells were pooled and propagated with immobilized anti-CD3 and anti-CD28, anti-CD4 was added to protect uninfected cells. All cells were plated at 2.5 x 10⁵ cells/well in 24-well plates. The number of CD4 + cells recovered after six days in culture is shown in Table 11.

Table 11 Pooled CD4+ Cells Purged of Active and Latent Virus Expanded 6 Days.

Day	Group 1	Group 2	Group 3
0	3.3 x 10 ⁶	2.1 x,10 ⁶	3.6 x 10 ⁶
6	12.4 x 10 ⁶	11.8 x 10 ⁶	11.4 x 10 ⁶ ;

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CD4+ cells purged of active and latent virus were expanded in 24-well plates. Cells were harvested and counted after six days in culture with immobilized anti-CD3 mAb-and anti-CD28 mAb.

The cells from the 24-well plates were pooled and incubated in spinner flasks for three days. They were then relabelled with anti-CD4 and rosetted with GAM-coated immunomagnetic beads. 1 x 10⁶ positively selected cells were co-cultured with indicator cells for 20 days. The cell-lysates for all three groups were negative for p24 (data not

5 shown). These results demonstrate that this method is capable of producing a viral-free fraction of CD4+ cells from the peripheral blood of AIDS patients.

The cells from the three groups were pooled and relabelled with anti-CD3 mAb and inoculated into 2 GAM-coated cartridges of a min20 hollow fiber device with 200 ng/ml of anti-CD28 mAb. After 21 days of culture, 1.7 x 10⁸ cells were harvested. Three days after harvest, the cells were relabelled with anti-CD3 mAb and inoculated into a single GAM-coated cartridge on the large scale device with 200 ng/ml of anti-CD28 mAb. After 21 days of culture, 1.1 x 10¹⁰ cells were harvested.
25 Three days after harvest, these cells were relabelled with anti-CD3 mAb and inoculated into 8 GAM-coated cartridges on the large-scale device with 200 ng/ml of anti-CD28 mAb. After 18 days of culture, 6.4 x 10¹⁰ CD4+ cells were recovered. The cells were negative for p24.

To demonstrate that CD4 + cells isolated and propagated by this process were still capable of normal function, their ability to enhance NK

CD4 + Functional Studies

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activity was assessed. Patients with AIDS are known to have reduced NK function. Some reports have shown that exogenous IL-2 can significantly enhance NK-function of AIDS patients in vitro. This study demonstrated that adding the expanded viral-purged CD4+ cells was effective.

Materials and methods

The NK-sensitive cell line K562 was used as the target cell. The cells were chromium labelled by suspension at a concentration of 1.cx.107 cells/ml, in cRPMI containing 1,00 pCl/ml-of (%Cr) sodium chromate (New England Nuclear, Boston, MA) for 60 minutes: at 37°C a The cells were then washed twice, resuspended at 5 x 10° cells/ml in 1,00 pl aliquots into wells of round-bottomed 96 well, plates.

Monocyte depleted lymphocytes from AIDS patients suspended at 5 x 10⁸ cells/ml were added to wells containing the target cells in 50 μ l aliquots. An additional 50 μ l of medium or CD4+ cells was added to each well such that the effector:target ratio without CD4+ cells was 50:1...

After a one hour incubation at 379°C in five percent CO₂ at 100 percent humidity, the plates were centrifuged at 800 × g for 12 minutes and 100 µl aliquots of each well were harvested and counted on a liquid scintillation counter. Percent lysis of each target cell was determined by the equation:

% lysis = cpm_{test} - cpm_{control} cpm_{max} - cpm_{control} x 100, where cpm_{test} indicates chromium counts per minute released in the presence of lymphocytes, cpm_{control} indicates release of the presence of medium alone, and cpm_{max} indicates release in the presence of BRIS-35 detergent (Sigma, St. Louis, MO).

Each test was performed in quadruplicate. Significance of percent lysis was determined by comparing mean cpm_{test} with mean cpm_{colivel} by student's t-test. Results are shown in Table 12.

Table 12. NK-Activity of Lymphocytes from AIDS Patient Supplemented with Autologous, Viral-Purged CD4+ Cells.

Results	% Lysis
AIDS lymphocytes alone	26.2 ± 6.5%
AIDS lymphocytes +1 IL-2 (10 U/ml)	54.5 ± 6,8%
AIDS lymphocytes + CD4+ (1000)	33.4 ± 7.0%
AIDS lymphocytes + CD4+ (5000)	48.8 ± 3.5%
AIDS lymphocytes + CD4+ (10,000)	64.6 ± 5%
AIDS lymphocytes + CD4+ (50,000)	64.2 ± 9.5%
Normal lymphocytes alone	60.2 ± 6.4%
Normal lymphocytes + IL-2 (10 U/ml)	73.5 ± 6.5%

NK-activity of a single AIDS patient after reconstruction with autologous/viralpurged CD4+ cells. The number of added cells is noted in parentheses. Results are expressed, as the magn. ASE of quadruplicate samples. 1

The NK-activity, of AIDS patients of 26.2 ± 6.5% was significantly lower than the 60.2 ± 6.4% for normal controls. The addition of IL-2 significantly increased NK-activity, in normal and AIDS patients, but had a much greater affect in AIDS. The addition of 1,000 autologous CD4+ cells did not significantly increase NK-activity. Addition of 5,000 and 10,000 CD4+ cells significantly increased activity to normal levels.

Addition of 50,000 CD4+ had the same effect as 10,000 cells

These results evidence that the CD4+ cells isolated and expanded by this protocol are able to produce IL-2. These results also support the evidence that large numbers of these CD4+ cells infused back to the patient should restore immunological function.

Purification of HIV-Specific T-cells

HIV-specific class I-restricted T-cells are known to be present in the blood of AIDS patients; they are presumed to be a subset of CD8+, CD28+, CD 11', CD25+ lymphocytes. These are in vivo activated (CD25+ same as IL2R+) Tc (CD28+ same as 9.3). To isolate these

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cells, a series of positive selection steps were conducted using CD8 (leu 2a, Becton Dickinson), CD28 (KOLT-2 gift from K. Sagawa), and CD25 (IL-2R, Coulter) mAbs and GAM-coated immunomagnetic beads.

Positive selection occurred in the following order: CD8, CD28, and finally, CD25. A subset of the isolated cells should be HIV-specific: The other in vivo T-cells in this group may also be of the apeutic importance; they may be specific for other adventitious agents afflicting the patient.

AIDS patients usually had a high percentage of CD25+ cells. In six patients tested, the mean CD25+ cells were 14 ± 8% compared to six normal controls at 3' ± 2.5%.

CD8+ Functional Studies

The CD8+ CD28+ CD25+ T-cells isolated from an AIDS patient and expanded to 5.3 x 10¹⁶ cells were tested for their ability to lyse HIV-infected autologous CD4+ lymphocytes. The target lymphocytes were expanded viral-free CD4+ cells from the same patient from whom the effector cells were isolated. The CD4+ cells were activated on immobilized anti-CD3 at 5 x 10⁶ cells/ml in ohe ml cRPMI on a 24-well plate. One ml of H9+ supernatant containing 10⁶ U/ml IL-2 was added to each well. The CD4+ cells were harvested from the wells after incubation at 37°C in five percent CO₂ at 100 percent humidity for four days.

The cells were labelled with ⁵¹Cr using the same procedure as described for K562 target cells. All cells were plated in round-bottomed 96-well plates at effector:target ratios of 100:1, 50:1, and 25:1. Percent lysis was determined as described earlier. Each test was performed in triplicate. Results are shown in Table 13.

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Table 13 CD8+, CD28+, CD25+ Killer T-Cells Isolated from HIV + $\sqrt{2}$ Patient, Ability to Lyse Autologous HIV Infected Cells

Cell: Target Ratio	% Lysis	in areally.
100:1	21.0 ± 8.0%	"Il via its proportionalism
	9.0 ± 3.5%	
2015 V (25:1/17026 B.C. V	3.5 ± 2.0%	mattern ore the

CD8+, CD28+, CD25+ To isolated from an AIDS patient were tested for their ability to lyse studiologious CD4+ cells infected with Hi-1. Percent lysis was calculated from a *Ucr-release assay.

These results indicate significant effector function. The low percentage lysis was probably due to a combination of a low percentage of targets infected with HIV (74 percent remained CD4+), and a high was background.

Although the present invention has been described with reference to preferred embodiments, workers skilled in the art will recognize that changes may be made in form and detail without departing from the spirit and scope of the invention. Since modifications will be apparent to those of skill in this art, it is intended that this invention be limited only by the scope of the appended claims.

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WHAT IS CLAIMED IS:

- A method for generating immune cells, comprising:
 collecting material comprising body fluid or tissue containing mononuclear cells from a mammal; and
- contacting, in the absence of exogenous interleukin-2, the material with one or more activating proteins specific for cell surface proteins present on cells in the material and in an amount sufficient to induce ex vivo cell expansion, whereby the cells expand to clinically relevant numbers.
 - 2. The method of claim 1, wherein prior to or during the contacting step, the cells in the material are treated under conditions whereby ex <u>vivo</u> differentiation of some or all of the cells into selected regulatory immune cells is induced.
- 3. The method of claim 1, wherein prior to or during the contacting step, the cells in the material are treated under conditions, whereby ex vivo differentiation of some or all of the cells into desired effector immune cells is induced.
 - 4. The method of any of claims 1-3, wherein the expanded cells are purified.
- The method of any of claims 1-4, wherein the immune cells are specific for a defined antigen.
 - 6. The method of any of claims 1-5, wherein the expanded cells are predominantly Th1, Th2 or Th3 cells.
- The method of claim 1 or claim 2, wherein the immune cells
 are activated <u>ex vivo</u> in the presence of either or both interferon-y and IL to cause differentiation of Th1 cells.
 - 8. The method of claim 1 or claim 2, wherein the cells are activated in the presence of IL-4 with or without the presence of anti-

gamma interferon and/or anti-IL-12 to cause the differentiation of Th2 cells.

- 9. The method of any of claims 1-8, wherein the proteins specific for cell surface molecules are one or more monoclonal antibodies specific for immune cell surface proteins.
 - 10. The method of claim 9, wherein the monoclonal antibodies are specific for CD3 or CD2, combined with any combination of one or more of the following: CD4, CD8, CD11a, CD27, CD28, CD44 and according to the combined control of the combined contr
- 10. The method of any of claim 1-10, wherein expansion is effected in a hollow fiber bioreactor.
 - .12. The method of any of claims 1-11, wherein the immune cells are expanded to an excess of 10° cells.
- .13. ::The method of any of claims 1-11, wherein the immune cells
 - 14. The method of claim 1 or claim 2, wherein the cells are effector immune cells:
 - 15... The method of claim 1-or claim 2, wherein the cells are regulatory immune cells.
 - 16... A method for autologous cell therapy; comprising:

 collecting material comprising body fluid or tissue containing:

 mononuclear cells from a mammal; and
- contacting, in the absence of exogenous interleukin-2, the material with one or more activating proteins specific for cell surface proteins:

 25 present on cells in the material and in an amount sufficient to induce ex vivo cell expansion, whereby the cells expand to clinically relevant to numbers; and

infusing the resulting cells into a mammal.

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- 17: The method of claim 16, wherein expanded cells are purified prior to infusion into the mammal.
- 18. paThe method of claim 16 or claim 17, Wherein the expanded cells are regulatory immune cells.
- 5 19. The method of claim 16 or claim 17, wherein the expanded cells are effector immune cells.
 - comprising: As the state of the

collecting material containing mononuclear cells from a mammal;
treating the cells to alter their cytokine production profile; and
expanding the cells to a clinically relevant number of cells.

- 21. The method of claim 20, wherein the immune cells with
- 22a a The method of claim 20, wherein the immune cells with
 - 23. The method of any of claims 20-22, wherein the mononuclear cells are treated to differentiate into Th1 or Th2 cells.
 - 24. The method of any of claims 20-22, wherein the resulting population of cells are Th1-like or Th2-like cells.
 - 25. The method of any of claims 20-22, wherein the immune cells are activated <u>ax vivo</u> in the presence of either or both interferon-y and IL-2 to cause differentiation of Th1 cells.
- 6.26. a:The method of claim:25, wherein anti-IL-4 mAb is also present during activation.
 - 27.c. The method of claim 26, wherein the effector cells are activated in the presence of IL-4 with or without the presence of anti-gamma interferon and/or anti-IL-12 antibodies to cause the differentiation of Th2 cells.

- The method of any of claims 20-27, wherein one or more monoclonal antibodies are included in the medium in which the mononuclear cells are expanded.
- 29. The method of claim 28, wherein the monoclonal antibodies are specific for CD3 or CD2, combined with any combination of one or more of the following: CD4, CD8, CD11a, CD27, CD28, CD44 and CD458O.
 - 30. The method of any of claims 20-29, wherein the cells are expanded in a hollow fiber bioreactor.
- 10 31. The method of any of claims 20-30, wherein the cells are expanded to an excess of 10° cells.
 - 32. The method of any of claims 20-30, wherein the cells are expanded to an excess of 10^{10} cells.
- 33. A method of producing virally purged CD4+ cells,
 - isolating CD4* cells from a patient infected with human immunodeficiency virus (HIV);

contacting the cells with one or more protein activating agents:

agents;

- selecting cells CD4* that are HIV; and then expanding the selected cells to clinically relevant numbers.
- 34. The method of claim 33, wherein in the contacting step, the activating under conditions promote Th1 cell differentiation.
- 25 35. The method of claim 33, further comprising:
 - after selecting CD4⁺ that are HIV and prior to expanding the selected cells, growing a plurality of aliquots in the presence of mitogenic agents:

selecting from the aliquots those that are HIV: and

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then expanding the selected cells to clinically relevant numbers.

- 36. The method of claim 33, wherein the cells are activated with anti-CD3 mAb in the presence of interferon (IFN-y).
- 37. The method of claim 35, wherein, after activation, the cells are grown in the presence of anti-CD28 mAb and IFN-V.
 - 38. The method of claim 1, wherein the cells are CD8+ cells.
 - A composition, comprising a clinically relevant number of CD4+ cells.
- 40. A composition comprising virally purged CD4* cells produced by the method of claim 33.
 - 41. The composition of claim 39, wherein the CD4+ cells are predominantly Th1-cells.
- The composition of claim 39, wherein the CD4+ cells are
 predominantly Th2-cells.
 - 43. A combination, comprising:

a composition containing a clinically relevant number of virelly-purged CD4+ cells; and

- a composition containing a clinically relevant number of 20 CD8* effector cells.
 - 44.7 A method of treating a patient infected with HIV, comprising:

administering a clinically relevant number of virally-purged CD4* cells.

25 45. The method of claim 44, further comprising administering a clinically relevant number of CD8* effector cells, wherein the effector cells are administered, before, after or simultaneously with the CD4* cells.

- $_{\rm comprising}$: A method of treating patients with autologous immune cells, comprising:
- collecting, a tissue or body fluid sample comprising mononuclear cells from a mammal; (a) and a property of the sample comprising mononuclear cells from a mammal;
- - reinfusing a sufficient number of the cells to alter the in vivo
- 47.... The method of claim 46, wherein the cells are treated to differentiate into Th2-like cells.
 - with an autoimmune disease or disease characterized by chronic inflammation.
- 49. The method of claim 46, wherein the cells are treated to the differentiate into Th1-like cells, and provide an area of the self.
 - 50. The method of claim 49, wherein the patients are diagnosed with allergic disorders or infectious disease.
 - "51. ... The method of claim 46, wherein the patients are to receive an organ or tissue transplant from an allogeneic or xenogeneic donor.
- 52₃. The method of claim 49, wherein the immune cells are exposed to one or more antigens from one or more pathogenic organisms and reinfused to protect the patient from subsequent infection; from the same pathogens, as a remain to you by market and the same pathogens.
- 5 human regulatory T-cells
 - 54. The composition of claim 53, wherein the cells are contained in a volume of one liter or less, present or contains from a to present the cells.
 - 55. The composition of claim 53, wherein the cells are to be a contained in a volume of 500 mls or less.

- The composition of claim 55, wherein the volume is 250 mls or less.
- 57. The composition of any of claims 53-56, wherein the concentration of cells is at least about 10⁷-10⁸ cells/mil. The concentration of cells is at least about 10⁷-10⁸ cells/mil.
- 58.45 The composition of any of claims 53-567 comprising at least 10° regulatory immune cells, now any or tensor by a drop immune cells, now any or tensor by a drop immune cells.
- 59::: A composition of claim-53; comprising at least 10° Th3 cells.
- 60.1 *The composition of claim-58 of claim-59, comprising at least 10 1010 cells.
 - the 1.61.1 The composition of any of claims 53-56, wherein the cells are Th1 cells. The composition of any of the production of the cells.
 - 62. The composition of any of claims 53-58, wherein the cells of are Th2tcells, the composition of companies are that the taxon and the cells of the composition of t
- 15 63. Use of the composition of any of claims 55 59 for the manufacture of a medicament for treating autoimmune disease.
 - 64. The use of the composition of any of claims 53:58; Wherein the disease is selected from rheumatoid arthritis, inflammatory bowel disease (IBD) or to prevent transplant rejection.
- 20 653 (*Use of claim 64) wherein the compositions is used for preventing rejection of transplanted islets for treatment of insulin because dependent diabetes mellitus.
 - 66. Use of the composition of any of claims 53-58 for the manufacture of a medicament for treating allergies, infectious disorders or diseases, tumors or as a vaccine.
 - hash 67... Use of the composition of any of claim 59 for the selfmanufacture of a medicament for treating multiple sclerosis or insulin-

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68.	a composition of any of claims 53-62; and a composition comprising a clinically relevant number of man effector T-cells. 69. The combination of claim 68, wherein the concentrations of man regulatory cells and human effector cells are each at least about 7-10° cells/ml. 70. The combination of claim 68 or claim 69, wherein the mpositions are mixed. 71. A method for treating autoimmune disorders, comprising ministering a composition containing a therapeutically effective number regulatory immune cells, whereby the symptoms of the disease are relicrated or progression of the disease is retarded. 72. The method of claim 71; wherein the disease is rheumatoid hritis, multiple sclerosis, insulin-dependent diabetes mellitus; on alarmatory bowel disease. 73. The method of claim 71, wherein the population of immune ls is Th2-like. 74. The method of claim 71, wherein the cells are contained in a ume of 1 liter or less. 75. The method of claim 74, wherein the disease is rheumatoid hritis, wherein the composition is produced by a method comprising: collecting mononuclear cells from a rheumatoid arthritis patient; expanding the cells under conditions whereby a composition intaining an amount of Th2 cells sufficient to suppress or reduce the onic inflammatory lesions of the arthritis; and		
	a composition comprising a clinic	ally relevant numb	er of
human effec	ctor T-cells.	A. 10, 11 . Late	ta anno 1
69.	The combination of claim 68, wh	erein the concentra	ations of
7,1.	A method for treating autoimmune	disorders, compris	ing
inflammator	y bowel disease.	s delivioloti Jeni pali.	L181
73.	The method of claim 71, wherein	the population of i	mmune
74.	The method of claim 71, wherein	the number of reg	ulatory
75.	The method of claim 74, wherein	the cells are conta	ined in a
76	The method of claim 71, wherein	the disease is rheu	matoid
collect	ting mononuclear cells from a rheu	matoid arthritis pat	tient;
expan	ding the cells under conditions wh	ereby a compositio	north and the
containing ar	n amount of Th2 cells sufficient to	suppress or reduc	e the

infusing the resulting composition of cells into the patient.

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- 77. The method of claim 76, wherein the number Th2 cells is at Sangara and Sangara and Sangara least 109.
- 78. The method of claim 76, wherein the cells are contained in a en en a voor marken? volume of 1 liter or less.
- 79. The method of claim 76, wherein the Th2 cells are memory cells, and the property of the property of the control of the property of the control of the con
 - The method of claim 79, wherein the Th2 cells are activated ex vivo in the presence of interferon-v. IL-2, or mixtures thereof, prior to Locality of the state of the infusion.
- 81. The method of claim 71, wherein the disease is multiple sclerosis, and the composition is produced by a method, comprising: collecting mononuclear cells from a multiple sclerosis patient: expanding the cells under conditions whereby a composition containing an amount of Th3 cells sufficient to ameliorate the symptoms 15 or retard or stop the progression of multiple sclerosis: and

infusing the resulting composition of cells into the patient. 82. The method of any of claims 71-81, wherein the number of cells is at least 109 cells.

- 83. The method of any of claims 71-82, wherein the cells are Carry Bayler II we obtain the second 20 contained in a volume of 1 liter or less.
 - 84. The method of any of claims 71-83, wherein the cells have a memory phenotype.
 - 85. The method of claim 81, wherein the cells are specific for myelin or encephalitogenic epitopes of myelin antigens.
 - 86. The method of claim 71, wherein the disease inflammatory bowel disease (IBD), and the composition is produced by a method. comprising: and a management of the constraint o

collecting mononuclear cells from an IBD patient:

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expanding the cells under conditions whereby a composition.

containing an amount of Th2 cells sufficient to ameliorate the symptoms or retard or stop the progression of the IBD; and the condition of cells into the patients.

- 87. The method of claim 86, wherein the number of cells is at least 10° cells.
- 88. The method of claim 86, wherein the cells are contained in a volume of 1 litter or less may appear to a dispression. The behave and granded
- 89. The method of claim 86, wherein the disease is Crohn's disease (CD) or ulcerative colitis (UC).
 - 90. The method of claim 86, wherein the Th2 cells are express integrin, a4, β7.
- 91. A method for suppression transplant rejection, comprising: collecting mononuclear cells from a patient prior to undergoing organ or tissue transplantation;

expanding the cells under conditions whereby a composition containing an amount of Th2 cells sufficient to prevent rejection of the transplanted organ or tissue; and

infusing the resulting composition of cells into the patient.

- 20 92. The method of claim 91, wherein the number of cells is at least 10° cells.
 - 93. The method of claim 91, wherein the cells are contained in a volume of 1 liter or less. $y_{ij} = y_{ij} \cos x_{ij} \cos x_{ij} \cos x_{ij}$
 - 94. The method of claim 91, wherein the transplanted tissue are transplanted islets of Langerhans.
 - 95. The method of claim 91, wherein the cells are specific for the alloantigens or for an antigen unique to the transplanted tissue or organ.

96. A method for treating insulin-dependent diabetes mellitus (IDDM), comprising:

collecting mononuclear cells from a patient diagnosed with IDDM or at high risk for developing IDDM;

expanding the cells under conditions whereby a composition containing an amount of Th2 cells sufficient to prevent or retard islet destruction; and

infusing the resulting composition of cells into the patient.

- 97. The method of claim 96, wherein the number of cells is at
 - 98. The method of claim 96 or claim 97, wherein the cells are contained in a volume of 1 liter or less.
 - 99. A method for treating allergies, comprising collecting mononuclear cells from a patient prior to undergoing organ or tissue transplantation:

expanding the cells under conditions whereby a composition containing an number of Th1 cells sufficient to ameliorate the symptoms of the allergy; and

infusing the resulting composition of cells into the patient.

- 100. The method of claim 99, wherein the number of cells is at least 10° cells.
- 101. The method of claim 99 or claim 100, wherein the cells are contained in a volume of 1 liter or less.
- 102. The method of any of claims 99-101, wherein the cells are specific for one or more allergens.
- 103. A method for treating infectious diseases or cancers, comprising:

collecting mononuclear cells from a patient prior to undergoing organ or tissue transplantation;

-89-

expanding the cells under conditions whereby a composition:

containing a therapeutically effective number of Th1 cells; and a containing a therapeutically effective number of Th1 cells; and a containing a contain

infusing the resulting composition of cells into the patient.

104. The method of claim 103, wherein the number of cells is at

- 105. The method of claim 103 or claim 104, wherein the cells are of contained in a volume of 1 liter, or less.
- 106. A method for treating infectious diseases or cancers, comprising: co-infusing therapeutically effective numbers of regulatory and effector cells.
 - 107. The method of claim 106, further comprising co-infusing CD8* effector cells cytotoxic T lymphocytes (CTLs) that are specific for the pathogen or tumor.
- 108. The method of claim, 106 or claim 107, wherein the
 - 109. The method of any of claims 106-108, wherein the regulatory cells are specific for the pathogen or tumor.
 - 110. The method of claim 103 or claim 106, wherein the disease is renal cell carcinoma and the antigen is Hsp70.
- 20 111. The method of any of claims 103-110, wherein the number of cells is at least 109 cells.
 - 112. The method of any of claims 103-111, wherein the cells are contained in a volume of 1 liter or less.
 - 113. A method of vaccination, comprising

exposing isolated mononuclear cells obtained from a patient to a selected vaccine antigen in the presence of one or more cytokines that induce Th1 cells or Th1-like cells to produce Th1 cells or Th1-like cells specific for the antigen; and

expanding the resulting cells for reinfusion.

- 114. The method of claim 113, wherein the number of cells is at least 10⁹ cells.
- 115. The method of claim 113 or claim 114, wherein the cells are contained in a volume of 1 liter of less.
- 5 116. The method of any of claims 113-115, wherein the cells have a memory phenotype.
 - 117. The method of any of claims 113-116, wherein the cytokine(s) is (are) selected from IL-12 and IFN-y.
- 118: The method of any of claims 113-115, wherein the resulting 10 cells are CD4+, CD8+ or a mixture thereof.
 - 119. A method for altering the regulatory balance of immune cells in a human, comprising administering to the human a composition comprising a clinically relevant number of autologous regulatory T-cells.
- 120. The method of claim 16, wherein at least 10° cells are
 - 121. The method of claim 16, wherein at least 1010 cells are administered.
 - 122. The method of claim 1 or claim 16, wherein the cells are Th1 cells.
- 20 123. The method of claim 1 or claim 16, wherein the cells are Th2 cells.
 - 123. The method of claim 1 or claim 15, wherein the cells are Th3 cells.
- 125. The method of claim 16, further comprising, administering a 25 clinically relevant number of effector immune cells, wherein the effector immune cells are administered with, before or after administration of the regulatory cells.
 - 126. Use of the composition of claim 53 for treating autoimmune disease

- 127. The use of claim 126, wherein the disease is selected from rheumatoid arthritis, inflammatory bowel disease (IBD) or to prevent transplant rejection.
- 128. Use of claim 126, wherein the compositions is used for . 5 preventing rejection of transplanted islets for treatment of insulin-dependent diabetes mellitus.
 - 129. Use of the composition of claim 53 for treating allergies, infectious disorders or diseases, turnors or as a vaccine.
- 130. Use of the composition of claim 59 for treating multiple sclerosis or insulin-dependent diabetes mellitus.
 - 131. The use of any of claims 126-130, wherein the composition contains at least 10° regulatory cells.
- 132. The use of any of claims 126-131, wherein the composition has a volume of 1 liter or less, preferably 500 mls or less, more preferably 250 mls or less.
 - 133. The use of claim 126, wherein the disease is rheumatoid arthritis.
 - The use of claim 126, wherein the disease is multiple sclerosis.
- 20 135. Use of the composition of any of claims 39-42 for treating HIV infection.
 - 136. Use of the composition of any of claims 39-42 for formulating a medicament for treatment of HIV infection.

INTERNATIONAL SEARCH REPORT

Tational Application No

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Ir *ational Application No PUT/US 96/12170

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International application No.

INTERNATIONAL SEARCH REPORT

PCT/US 96/12170

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
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1. X	Claims Nos
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ب ٠	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the charges; it is convered by claims Nos:
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INTERNATIONAL SEARCH REPORT

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/210

PCT/ISA/210

Remark: Although claims 16-19, 44-52, 71-112, 119-121, 125-135 completely, and 122, 123 partially, are directed to a method of treatment of the human/animal body the search has been carried out and based on the alleged effects of the compound/composition.

INTERNATIONAL SEARCH REPORT

Information on patent family members

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